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(54) Title: NOVEL HUMAN GENES AND GENE EXPRESSION PRODUCTS

(57) Abstract: The invention provides novel polynucleotides. The invention further provides novel members of protein families, and polynucleotides that are differentially expressed in cancer cells relative to normal cells, and in metastatic cancer cells relative to normal cells or non-metastatic cancer cells.

NOVEL HUMAN GENES AND GENE EXPRESSION PRODUCTS

FIELD OF THE INVENTION

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The present invention relates to novel polynucleotides of human origin and the encoded gene products.

5 BACKGROUND OF THE INVENTION

Identification of novel polynucleotides, particularly those that encode an expressed gene product, is important in the advancement of drug discovery, diagnostic technologies, and the understanding of the progression and nature of complex diseases such as cancer. Identification of genes expressed in different cell types isolated from sources that differ in disease state or stage, developmental stage, exposure to various environmental factors, the tissue of origin, the species from which the tissue was isolated, and the like is key to identifying the genetic factors that are responsible for the phenotypes associated with these various differences.

This invention provides novel human polynucleotides, the polypeptides encoded by these polynucleotides, and the genes and proteins corresponding to these novel polynucleotides.

SUMMARY OF THE INVENTION

This invention relates to novel human polynucleotides and variants thereof, their encoded polypeptides and variants thereof, to genes corresponding to these polynucleotides and to proteins expressed by the genes. The invention also relates to diagnostics and therapeutics comprising such novel human polynucleotides, their corresponding genes or gene products, including probes, antisense nucleotides, and antibodies. The polynucleotides of the invention correspond to a polynucleotide comprising the sequence information of at least one of SEQ ID NOs: 1-3351.

Various aspects and embodiments of the invention will be readily apparent to the ordinarily skilled artisan upon reading the description provided herein.

DETAILED DESCRIPTION OF THE INVENTION

The invention relates to polynucleotides comprising the disclosed nucleotide sequences, to full length cDNA, mRNA genomic sequences, and genes

corresponding to these sequences and degenerate variants thereof, and to polypeptides encoded by the polynucleotides of the invention and polypeptide variants.

Polypeptide variants differ from wild type protein in having one or more amino acid substitutions that either enhance, add, or diminish a biological activity of the wild type protein.

Six of the polypeptides disclosed herein encode new members of the MKK kinase family; the coding region is found within the nucleotide region in parentheses: SEQ ID NO:29 (nucleotides 295-421); SEQ ID NO:31 (298-397); SEQ ID NO:196 (37-322); SEQ ID NO:3175 (nucleotides 14-164); SEQ ID NO:3190 (229-390); and SEQ ID NO:3281 (15-182). Twenty-four of the polypeptides encode new members of the family of transcription factor proteins having a basic region plus leucine zipper: SEQ ID NO:410 (42-191); SEQ ID NO:552 (116-288); SEQ ID NO:768 (116-288); SEQ ID NO:822 (108-262); SEQ ID NO:836 (158-353); SEQ ID NO:1288 (73-234); SEQ ID NO:1365 (69-257); SEQ ID NO:1540 (289-471); SEQ ID NO:1549 (200-391); SEQ ID NO:1556 (163-354); SEQ ID NO:1557 (207-398); SEQ ID NO:1563 (107-298); SEQ ID NO:1622 (180-365); SEQ ID NO:1630 (100-291); SEQ ID NO:1704 (184-372); SEQ ID NO:1808 (36-161); SEQ ID NO:1454 (49-209); SEQ ID NO:2363 (48-211); SEQ ID NO:2424 (43-194); SEQ ID NO:3147 (190-369); SEQ ID NO:3152 (129-320); SEQ ID NO:3158 (167-334); and SEQ ID NO:3208 (34-256).

SEQ 1D NOs:186 (175-395); 2591 (60-165); 3307 (43-321); and 3339 (94-342) encode polypeptides having an SH2 domain, and SEQ 1D NOs:234 (23-121), 1832 (18-173), and 1835 (57-206) encode polypeptides having an SH3 domain. Nine polypeptides encode new members of the family of proteins having Ank repeat regions: SEQ 1D NO:187 (358-432); SEQ 1D NO:1268 (238-315); SEQ ID NO:1804 (301-378); SEQ ID NO:1819 (278-355); SEQ ID NO:1839 (224-307); SEQ ID NO:1830 (184-267); SEQ ID NO:2562 (18-101); SEQ ID NO:3015 (131-214); and SEQ ID NO:3267 (97-180).

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The following eleven polynucleotides encode polypeptides having a C2H2 type zinc finger: SEQ ID NOs:308 (110-172); 807 (339-392); 1324 (294-356); 1503 (154-216); 1527 (156-212); 1674 (196-258); 1779 (64-126); 1801 (295-351); 3081 (190-252); 3193 (293-355); and 3306 (161-223). Eight polynucleotides encode polypeptides of the family of ATPases: SEQ ID NOs:431 (71-428); 639 (157-561); 2135 (2-401); 2684 (9-461); 2859 (100-320); 3178 (45-386); 3197 (281-343) and 3266 (8-139). Polypeptides having a fibronectin type III domain are encoded by SEQ ID NO:746 (209-427) and 1192 (186-416). Polypeptides having an EF-hand domain are encoded by SEQ ID NO:820 (341-

406); 1755 (281-367) and 3285(16-102). Six polypeptides of the protein kinase family are encoded by SEQ ID NOs:1157 (41-444); 1478 (54-437), 1496 (241-520); 2286 (12-182); 2969 (5-387); and 3190 (118-390).

LIM domain-containing polypeptides are encoded by SEQ ID NO:1269 (79-240); 1309 (248-404); 1360 (222-377); and 1386 (243-398). Two polypeptides of the family having a C2 domain (protein kinase C-like) are encoded by SEQ ID NO:1325 (1-234) and 2282(183-353). Polypeptides having a WD domain, G-beta repeat motif are encoded by SEQ ID NOs:1336 (66-164); 1380 (42-140); 1711 (263-361); 1762 (236-334); 1909 (160-258); 2218 (127-225); 3047 (191-292); 3108 (275-367) and 3292 (208-300).

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SEQ ID NO:1410 (222-350) encodes a member of the trypsin family. SEQ ID NOs:1417 (8-354); 2281 (20-387) and 2310 (20-371) encode members of the protein tyrosine phosphatase family. SEQ ID NOs:1464 (4-180) and 1514 (2-252) encode members of the family having an RNA recognition motif (also known as RRM, RBD, or RNP domain). SEQ ID NOs:1496 (241-520) and 3297(7-153) encode helicases having a conserved C-terminal domain. SEQ ID NO:1538 (9-635) encodes a member of the wnt family of developmental signaling proteins.

Three polynucleotides encode polypeptides having a homeobox domain: SEQ ID NOs:1676 (9-86); 1820 (123-299); and 1821 (127-303). A novel thioredoxin is encoded by SEQ ID NO:1677 (316-369). Two novel members of the ras family are encoded by SEQ ID NO:1688(109-410) and 3258(138-394). A novel polypeptide having a phosphatidylinositol-specific phospholipase C Y-domain is encoded by SEQ ID NO:1707 (92-439). A novel serine carboxypeptidase is encoded by SEQ ID NO:1744 (238-433). A novel polypeptide having N-terminal homology in the Ets domain is encoded by SEQ ID NO:1811 (184-315). A novel polypeptide having a bromodomain is encoded by SEQ ID NO:1814 (127-294). A novel polypeptide having a double-stranded RNA binding motif is encoded by SEQ ID NO:1818 (9-146). A novel polypeptide having a G-protein alpha subunit is encoded by SEQ ID NO:1846 (12-398).

SEQ ID NOs:1911 (35-151) and 1980 (60-197) encode polypeptides having a C3HC4 type zinc finger domain (RING finger). SEQ ID NO:2065 (253-306) encodes a polypeptide having a CCHC zinc finger domain. SEQ ID NO:2216 (90-179) encodes a polypeptide having a WW/rsp5/WWP domain. SEQ ID NO:2428 (25-350) encodes a polypeptide member of the dual specificity phosphatase family, having a catalytic domain.

SEQ ID NOs:2577 (0-311); 3183 (14-215); and 3195 (0-215) encode members of the 4 transmembrane segment integral membrane protein family. SEQ ID

NOs:2826 (116-400) and 2871 (198-392) encode polypeptides of the DEAD and DEAH box helicase family. SEQ ID NO:2944 (18-281) encodes a polypeptide having a calpain large subunit, domain III.

SEQ ID NO:3274 (11-187) encodes a eukaryotic transcription factor with a fork head domain. SEQ ID NO:3345 (65-271) encodes a polypeptide having a PDZ domain, and SEQ ID NO:3351 (124-270) encodes a polypeptide in the family of phorbol esters/glycerol binding proteins.

Described below are polynucleotide compositions encompassed by the invention, methods for obtaining cDNA or genomic DNA encoding a full-length gene product, expression of these polynucleotides and genes, identification of structural motifs of the polynucleotides and genes, identification of the function of a gene product encoded by a gene corresponding to a polynucleotide of the invention, use of the provided polynucleotides as probes and in mapping and in tissue profiling, use of the corresponding polypeptides and other gene products to raise antibodies, and use of the polynucleotides and their encoded gene products for therapeutic and diagnostic purposes.

Polynucleotide Compositions

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The scope of the invention with respect to polynucleotide compositions includes, but is not necessarily limited to, polynucleotides having a sequence set forth in any one of SEQ ID NOs:1-3351; polynucleotides obtained from the biological materials described herein or other biological sources (particularly human sources) by hybridization under stringent conditions (particularly conditions of high stringency); genes corresponding to the provided polynucleotides; variants of the provided polynucleotides and their corresponding genes, particularly those variants that retain a biological activity of the encoded gene product (e.g., a biological activity ascribed to a gene product corresponding to the provided polynucleotides as a result of the assignment of the gene product to a protein family(ies) and/or identification of a functional domain present in the gene product). Other nucleic acid compositions contemplated by and within the scope of the present invention will be readily apparent to one of ordinary skill in the art when provided with the disclosure here. "Polynucleotide" and "nucleic acid" as used herein with reference to nucleic acids of the composition is not intended to be limiting as to the length or structure of the nucleic acid unless specifically indicated.

The invention features polynucleotides that are expressed in human tissue, specifically human colon, breast, and/or lung tissue. Novel nucleic acid

compositions of the invention comprise a sequence set forth in any one of SEQ ID NOs:1-3351 or an identifying sequence thereof. An "identifying sequence" is a contiguous sequence of residues at least about 10 nt to about 20 nt in length, usually at least about 50 nt to about 100 nt in length, that uniquely identifies a polynucleotide sequence, e.g., exhibits less than 90%, usually less than about 80% to about 85% sequence identity to any contiguous nucleotide sequence of more than about 20 nt. Thus, the subject novel nucleic acid compositions include full length cDNAs or mRNAs that encompass an identifying sequence of contiguous nucleotides from any one of SEQ ID NOs:1-3351.

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The polynucleotides of the invention also include polynucleotides having sequence similarity or sequence identity. Nucleic acids having sequence similarity are detected by hybridization under low stringency conditions, for example, at 50°C and 10XSSC (0.9 M saline/0.09 M sodium citrate) and remain bound when subjected to washing at 55°C in 1XSSC. Sequence identity can be determined by hybridization under stringent conditions, for example, at 50°C or higher and 0.1XSSC (9 mM saline/0.9 mM sodium citrate). Hybridization methods and conditions are well known in the art, see, e.g., U.S. Patent No. 5,707,829. Nucleic acids that are substantially identical to the provided polynucleotide sequences, e.g., allelic variants, genetically altered versions of the gene, etc., bind to the provided polynucleotide sequences (SEQ ID NOs:1-3351) under stringent hybridization conditions. By using probes, particularly labeled probes of DNA sequences, one can isolate homologous or related genes. The source of homologous genes can be any species, e.g., primate species, particularly human; rodents, such as rats and mice; canines, felines, bovines, ovines, equines, yeast, nematodes, etc.

Preferably, hybridization is performed using at least 15 contiguous nucleotides (nt) of at least one of SEQ ID NOs:1-3351. That is, when at least 15 contiguous nt of one of the disclosed SEQ ID NOs. is used as a probe, the probe will preferentially hybridize with a nucleic acid comprising the complementary sequence, allowing the identification and retrieval of the nucleic acids that uniquely hybridize to the selected probe. Probes from more than one SEQ ID NO. can hybridize with the same nucleic acid if the cDNA from which they were derived corresponds to one mRNA. Probes of more than 15 nt can be used, e.g., probes of from about 18 nt to about 100 nt, but 15 nt represents sufficient sequence for unique identification.

The polynucleotides of the invention also include naturally occurring variants of the nucleotide sequences (e.g., degenerate variants, allelic variants).

Variants of the polynucleotides of the invention are identified by hybridization of putative variants with nucleotide sequences disclosed herein, preferably by hybridization under stringent conditions. For example, by using appropriate wash conditions, variants of the polynucleotides of the invention can be identified where the allelic variant exhibits at most about 25-30% base pair (bp) mismatches relative to the selected polynucleotide probe. In general, allelic variants contain 15-25% bp mismatches, and can contain as little as even 5-15%, or 2-5%, or 1-2% bp mismatches, as well as a single bp mismatch.

The invention also encompasses homologs corresponding to the polynucleotides of SEQ ID NOs:1-3351, where the source of homologous genes can be any mammalian species, e.g., primate species, particularly human; rodents, such as rats; canines, felines, bovines, ovines, equines, yeast, nematodes, etc. Between mammalian species, e.g., human and mouse, homologs generally have substantial sequence similarity, e.g., at least 75% sequence identity, usually at least 90%, more usually at least 95% between nucleotide sequences. Sequence similarity is calculated based on a reference sequence, which may be a subset of a larger sequence, such as a conserved motif, coding region, flanking region, etc. A reference sequence will usually be at least about 18 contiguous nt long, more usually at least about 30 nt long, and may extend to the complete sequence that is being compared. Algorithms for sequence analysis are known in the art, such as BLAST, described in Altschul et al., J. Mol. Biol. (1990) 215:403-10.

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In general, variants of the invention have a sequence identity greater than at least about 65%, preferably at least about 75%, more preferably at least about 85%, and can be greater than at least about 90%, 91%, 92%, 93%, 94%, 95%, or 96%, most preferably 97%, 98% or 99%. For the purposes of this invention, a preferred method of calculating percent identity is the Smith-Waterman algorithm, using the following. Global DNA sequence identity must be greater than 65% as determined by the Smith-Waterman homology search algorithm as implemented in MPSRCH program (Oxford Molecular) using an affine gap search with the following search parameters: gap open penalty, 12; and gap extension penalty, 1.

The subject nucleic acids can be cDNAs or genomic DNAs, as well as fragments thereof, particularly fragments that encode a biologically active gene product and/or are useful in the methods disclosed herein (e.g., in diagnosis, as a unique identifier of a differentially expressed gene of interest, etc.). The term "cDNA" as used herein is intended to include all nucleic acids that share the arrangement of sequence

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elements found in native mature mRNA species, where sequence elements are exons and 3' and 5' non-coding regions. Normally mRNA species have contiguous exons, with the intervening introns, when present, being removed by nuclear RNA splicing, to create a continuous open reading frame encoding a polypeptide of the invention.

A genomic sequence of interest comprises the nucleic acid present between the initiation codon and the stop codon, as defined in the listed sequences, including all of the introns that are normally present in a native chromosome. It can further include the 3' and 5' untranslated regions found in the mature mRNA. It can further include specific transcriptional and translational regulatory sequences, such as promoters, enhancers, etc., including about 1 kb, but possibly more, of flanking genomic DNA at either the 5' and 3' end of the transcribed region. The genomic DNA can be isolated as a fragment of 100 kbp or smaller; and substantially free of flanking chromosomal sequence. The genomic DNA flanking the coding region, either 3' and 5', or internal regulatory sequences as sometimes found in introns, contains sequences required for proper tissue, stage-specific, or disease-state specific expression.

The nucleic acid compositions of the subject invention can encode all or a part of the subject polypeptides. Double or single stranded fragments can be obtained from the DNA sequence by chemically synthesizing oligonucleotides in accordance with conventional methods, by restriction enzyme digestion, by PCR amplification, etc. Isolated polynucleotides and polynucleotide fragments of the invention comprise at least about 10, about 15, about 20, about 35, about 50, about 100, about 150 to about 200, about 250 to about 300, or about 350 contiguous nt selected from the polynucleotide sequences as shown in SEQ ID NOs:1-3351. The fragments also include those of lengths intermediate to the specifically mentioned lengths, such as 35, 36, 37, 38, 39, etc.; 150, 151, 152, 153, 154, etc. For the most part, fragments will be of at least 15 nt, usually at least 18 nt or 25 nt, and up to at least about 50 contiguous nt in length or more. In a preferred embodiment, the polynucleotide molecules comprise a contiguous sequence of at least 12 nt selected from the group consisting of the polynucleotides shown in SEQ ID NOs:1-3351.

Probes specific to the polynucleotides of the invention can be generated using the polynucleotide sequences disclosed in SEQ ID NOs:1-3351. The probes are preferably at least about a 12, 15, 16, 18, 20, 22, 24, or 25 nt fragment of a corresponding contiguous sequence of SEQ ID NOs:1-3351, and can be less than 2, 1, 0.5, 0.1, or 0.05 kb in length. The probes can be synthesized chemically or can be generated from longer polynucleotides using restriction enzymes. The probes can be

labeled, for example, with a radioactive, biotinylated, or fluorescent tag. Preferably, probes are designed based upon an identifying sequence of a polynucleotide of one of SEQ ID NOs:1-3351. More preferably, probes are designed based on a contiguous sequence of one of the subject polynucleotides that remain unmasked following application of a masking program for masking low complexity (e.g., XBLAST) to the sequence., i.e., one would select an unmasked region, as indicated by the polynucleotides outside the poly-n stretches of the masked sequence produced by the masking program.

The polynucleotides of the subject invention are isolated and obtained in substantial purity, generally as other than an intact chromosome. Usually, the polynucleotides, either as DNA or RNA, will be obtained substantially free of other naturally-occurring nucleic acid sequences, generally being at least about 50%, usually at least about 90% pure and are typically "recombinant", e.g., flanked by one or more nucleotides with which it is not normally associated on a naturally occurring chromosome.

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The polynucleotides of the invention can be provided as a linear molecule or within a circular molecule, and can be provided within autonomously replicating molecules (vectors) or within molecules without replication sequences. Expression of the polynucleotides can be regulated by their own or by other regulatory sequences known in the art. The polynucleotides of the invention can be introduced into suitable host cells using a variety of techniques available in the art, such as transferrin polycation-mediated DNA transfer, transfection with naked or encapsulated nucleic acids, liposome-mediated DNA transfer, intracellular transportation of DNA-coated latex beads, protoplast fusion, viral infection, electroporation, gene gun, calcium phosphate-mediated transfection, and the like.

The subject nucleic acid compositions can be used to, for example, produce polypeptides, as probes for the detection of mRNA of the invention in biological samples (e.g., extracts of human cells) to generate additional copies of the polynucleotides, to generate ribozymes or antisense oligonucleotides, and as single stranded DNA probes or as triple-strand forming oligonucleotides. The probes described herein can be used to, for example, determine the presence or absence of the polynucleotide sequences as shown in SEQ ID NOs:1-3351 or variants thereof in a sample. These and other uses are described in more detail below.

Use of Polynucleotides to Obtain Full-Length cDNA, Gene, and Promoter Region

Full-length cDNA molecules comprising the disclosed polynucleotides are obtained as follows. A polynucleotide having a sequence of one of SEQ ID NOs:1-3351, or a portion thereof comprising at least 12, 15, 18, or 20 nt, is used as a hybridization probe to detect hybridizing members of a cDNA library using probe design methods, cloning methods, and clone selection techniques such as those described in U.S. Patent No. 5,654,173. Libraries of cDNA are made from selected tissues, such as normal or tumor tissue, or from tissues of a mammal treated with, for example, a pharmaceutical agent. Preferably, the tissue is the same as the tissue from which the polynucleotides of the invention were isolated, as both the polynucleotides described herein and the cDNA represent expressed genes. Most preferably, the cDNA library is made from the biological material described herein in the Examples. The choice of cell type for library construction can be made after the identity of the protein encoded by the gene corresponding to the polynucleotide of the invention is known. This will indicate which tissue and cell types are likely to express the related gene, and thus represent a suitable source for the mRNA for generating the cDNA. As described in the Examples, cDNA of the invention was isolated from specific cell or tissue types, and such cells and tissues are preferable for obtaining related nucleic acids.

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Techniques for producing and probing nucleic acid sequence libraries are described, for example, in Sambrook et al., *Molecular Cloning: A Laboratory Manual, 2nd Ed.*, (1989) Cold Spring Harbor Press, Cold Spring Harbor, NY. The cDNA can be prepared by using primers based on sequence from SEQ ID NOs:1-3351. In one embodiment, the cDNA library can be made from only poly-adenylated mRNA. Thus, poly-T primers can be used to prepare cDNA from the mRNA.

Members of the library that are larger than the provided polynucleotides, and preferably that encompass the complete coding sequence of the native message, are obtained. In order to confirm that the entire cDNA has been obtained, RNA protection experiments are performed as follows. Hybridization of a full-length cDNA to an mRNA will protect the RNA from RNase degradation. If the cDNA is not full length, then the portions of the mRNA that are not hybridized will be subject to RNase degradation. This is assayed, as is known in the art, by changes in electrophoretic mobility on polyacrylamide gels, or by detection of released monoribonucleotides. Sambrook et al., *Molecular Cloning: A Laboratory Manual, 2nd Ed.*, (1989) Cold Spring Harbor Press, Cold Spring Harbor, NY. In order to obtain additional sequences

5' to the end of a partial cDNA, 5' RACE (PCR Protocols: A Guide to Methods and Applications, (1990) Academic Press, Inc.) can be performed.

Genomic DNA is isolated using the provided polynucleotides in a manner similar to the isolation of full-length cDNAs. Briefly, the provided polynucleotides, or portions thereof, are used as probes to libraries of genomic DNA. Preferably, the library is obtained from the cell type that was used to generate the polynucleotides of the invention, but this is not essential. Most preferably, the genomic DNA is obtained from the biological material described herein in the Examples. Such libraries can be in vectors suitable for carrying large segments of a genome, such as P1 or YAC, as described in detail in Sambrook et al., 9.4-9.30. In addition, genomic sequences can be isolated from human BAC libraries, which are commercially available from Research Genetics, Inc., Huntsville, Alabama, USA, for example. In order to obtain additional 5' or 3' sequences, chromosome walking is performed, as described in Sambrook et al., such that adjacent and overlapping fragments of genomic DNA are isolated. These are mapped and pieced together, as is known in the art, using restriction digestion enzymes and DNA ligase.

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Using the polynucleotide sequences of the invention, corresponding full-length genes can be isolated using both classical and PCR methods to construct and probe cDNA libraries. Using either method, Northern blots, preferably, are performed on a number of cell types to determine which cell lines express the gene of interest at the highest level. Classical methods of constructing cDNA libraries are taught in Sambrook et al., *supra*. With these methods, cDNA can be produced from mRNA and inserted into viral or expression vectors. Typically, libraries of mRNA comprising poly(A) tails can be produced with poly(T) primers. Similarly, cDNA libraries can be produced using the instant sequences as primers.

PCR methods are used to amplify the members of a cDNA library that comprise the desired insert. In this case, the desired insert will contain sequence from the full length cDNA that corresponds to the instant polynucleotides. Such PCR methods include gene trapping and RACE methods as described in Gruber et al., WO 95/04745 and Gruber et al., U.S. Patent No. 5,500,356. Kits are commercially available to perform gene trapping experiments from, for example, Life Technologies, Gaithersburg, Maryland, USA. In preferred embodiments of RACE, a common primer is designed to anneal to an arbitrary adaptor sequence ligated to cDNA ends (Apte and Siebert, *Biotechniques* (1993) 15:890-893; Edwards et al., *Nuc. Acids Res.* (1991) 19:5227-5232). When a single gene-specific RACE primer is paired with the common

primer, preferential amplification of sequences between the single gene specific primer and the common primer occurs. Commercial cDNA pools modified for use in RACE are available.

The promoter region of a gene generally is located 5' to the initiation site for RNA polymerase II. Hundreds of promoter regions contain the "TATA" box, a sequence such as TATTA or TATAA, which is sensitive to mutations. The promoter region can be obtained by performing 5' RACE using a primer from the coding region of the gene. Alternatively, the cDNA can be used as a probe for the genomic sequence, and the region 5' to the coding region is identified by "walking up." If the gene is highly expressed or differentially expressed, the promoter from the gene can be of usc in a regulatory construct for a heterologous gene.

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Once the full-length cDNA or gene is obtained, DNA encoding variants can be prepared by site-directed mutagenesis, described in detail in Sambrook et al., 15.3-15.63. The choice of codon or nucleotide to be replaced can be based on disclosure herein on optional changes in amino acids to achieve altered protein structure and/or function.

As an alternative method to obtaining DNA or RNA from a biological material, nucleic acid comprising nucleotides having the sequence of one or more polynucleotides of the invention can be synthesized. Thus, the invention encompasses nucleic acid molecules ranging in length from 15 nt (corresponding to at least 15 contiguous nt of one of SEQ ID NOs:1-3351) up to a maximum length suitable for one or more biological manipulations, including replication and expression, of the nucleic acid molecule. The invention includes but is not limited to (a) nucleic acid having the size of a full gene, and comprising at least one of SEQ ID NOs:1-3351; (b) the nucleic acid of (a) also comprising at least one additional polynucleotide or gene, operably linked to permit expression of a fusion protein; (c) an expression vector comprising (a) or (b); (d) a plasmid comprising (a) or (b); and (e) a recombinant viral particle comprising (a) or (b). Once provided with the polynucleotides disclosed herein, construction or preparation of (a) - (e) are well within the skill in the art.

The sequence of a nucleic acid comprising at least 15 contiguous nt of at least any one of SEQ ID NOs:1-3351, preferably the entire sequence of at least any one of SEQ ID NOs:1-3351, is not limited and can be any sequence of A, T, G, and/or C (for DNA) and A, U, G, and/or C (for RNA) or modified bases thereof, including inosine and pseudouridine. The choice of sequence will depend on the desired function and can be dictated by coding regions desired, the intron-like regions desired, and the

regulatory regions desired. Where the entire sequence of any one of SEQ ID NOs:1-3351 is within the nucleic acid, the nucleic acid obtained is referred to herein as a polynucleotide comprising the sequence of any one of SEQ ID NOs:1-3351.

Expression of Polypeptide Encoded by Full-Length cDNA or Full-Length Gene

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The provided polynucleotides (e.g., a polynucleotide having a sequence of one of SEQ ID NOs:1-3351), the corresponding cDNA, or the full-length gene is used to express a partial or complete gene product. Constructs of polynucleotides having sequences of SEQ ID NOs:1-3351 can be generated synthetically. Alternatively, single-step assembly of a gene and entire plasmid from large numbers of oligodeoxyribonucleotides is described by, e.g., Stemmer et al., Gene (Amsterdam) (1995) 164(1):49-53. In this method, assembly PCR (the synthesis of long DNA sequences from large numbers of oligodeoxyribonucleotides (oligos)) is described. The method is derived from DNA shuffling (Stemmer, Nature (1994) 370:389-391), and does not rely on DNA ligase, but instead relies on DNA polymerase to build increasingly longer DNA fragments during the assembly process.

Appropriate polynucleotide constructs are purified using standard recombinant DNA techniques as described in, for example, Sambrook et al., *Molecular Cloning: A Laboratory Manual, 2nd Ed.*, (1989) Cold Spring Harbor Press, Cold Spring Harbor, NY, and under current regulations described in United States Dept. of HHS, National Institute of Health (NIH) Guidelines for Recombinant DNA Research. The gene product encoded by a polynucleotide of the invention is expressed in any expression system, including, for example, bacterial, yeast, insect, amphibian and mammalian systems. Vectors, host cells and methods for obtaining expression in same are well known in the art. Suitable vectors and host cells are described in U.S. Patent No. 5,654,173.

Polynucleotide molecules comprising a polynucleotide sequence provided herein are generally propagated by placing the molecule in a vector. Viral and non-viral vectors are used, including plasmids. The choice of plasmid will depend on the type of cell in which propagation is desired and the purpose of propagation. Certain vectors are useful for amplifying and making large amounts of the desired DNA sequence. Other vectors are suitable for expression in cells in culture. Still other vectors are suitable for transfer and expression in cells in a whole animal or person. The choice of appropriate vector is well within the skill of the art. Many such vectors are

available commercially. Methods for preparation of vectors comprising a desired sequence are well known in the art.

The polynucleotides set forth in SEQ ID NOs:1-3351 or their corresponding full-length polynucleotides are linked to regulatory sequences as appropriate to obtain the desired expression properties. These can include promoters (attached either at the 5' end of the sense strand or at the 3' end of the antisense strand), enhancers, terminators, operators, repressors, and inducers. The promoters can be regulated or constitutive. In some situations it may be desirable to use conditionally active promoters, such as tissue-specific or developmental stage-specific promoters. These are linked to the desired nucleotide sequence using the techniques described above for linkage to vectors. Any techniques known in the art can be used.

When any appropriate host cells or organisms are used to replicate and/or express the polynucleotides or nucleic acids of the invention, the resulting replicated nucleic acid, RNA, expressed protein or polypeptide, is within the scope of the invention as a product of the host cell or organism. The product is recovered by any appropriate means known in the art.

Once the gene corresponding to a selected polynucleotide is identified, its expression can be regulated in the cell to which the gene is native. For example, an endogenous gene of a cell can be regulated by an exogenous regulatory sequence as disclosed in U.S. Patent No. 5,641,670.

Identification of Functional and Structural Motifs of Novel Genes

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Translations of the nucleotide sequence of the provided polynucleotides, cDNAs or full genes can be aligned with individual known sequences. Similarity with individual sequences can be used to determine the activity of the polypeptides encoded by the polynucleotides of the invention. Also, sequences exhibiting similarity with more than one individual sequence can exhibit activities that are characteristic of either or both individual sequences.

The full length sequences and fragments of the polynucleotide sequences of the nearest neighbors can be used as probes and primers to identify and isolate the full length sequence corresponding to provided polynucleotides. The nearest neighbors can indicate a tissue or cell type to be used to construct a library for the full-length sequences corresponding to the provided polynucleotides.

Typically, a selected polynucleotide is translated in all six frames to determine the best alignment with the individual sequences. The sequences disclosed

herein in the Sequence Listing are in a 5' to 3' orientation and translation in three frames can be sufficient. These amino acid sequences are referred to, generally, as query sequences, which will be aligned with the individual sequences. Databases with individual sequences are described in "Computer Methods for Macromolecular Sequence Analysis" Methods in Enzymology (1996) 266, Doolittle, Academic Press, Inc., a division of Harcourt Brace & Co., San Diego, California, USA. Databases include Genbank, EMBL, and DNA Database of Japan (DDBJ).

Query and individual sequences can be aligned using the methods and computer programs described above, and include BLAST, available over the world wide web at http://www.ncbi.nlm.nhi.gov/BLAST. Another alignment algorithm is Fasta, available in the Genetics Computing Group (GCG) package, Madison, Wisconsin, USA, a wholly owned subsidiary of Oxford Molecular Group, Inc. Other techniques for alignment are described in Doolittle, supra. Preferably, an alignment program that permits gaps in the sequence is utilized to align the sequences. The Smith-Waterman is one type of algorithm that permits gaps in sequence alignments. See Meth. Mol. Biol. (1997) 70: 173-187. Also, the GAP program using the Needleman and Wunsch alignment method can be utilized to align sequences. An alternative search strategy uses MPSRCH software, which runs on a MASPAR computer. MPSRCH uses a Smith-Waterman algorithm to score sequences on a massively parallel computer. This approach improves ability to identify sequences that are distantly related matches, and is especially tolerant of small gaps and nucleotide sequence errors. Amino acid sequences encoded by the provided polynucleotides can be used to search both protein and DNA databases.

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High Similarity. In general, in alignment results considered to be of high similarity, the percent of the alignment region length is typically at least about 55% of total length query sequence; more typically, at least about 58%; even more typically; at least about 60% of the total residue length of the query sequence. Usually, percent length of the alignment region can be as much as about 62%; more usually, as much as about 64%; even more usually, as much as about 66%. Further, for high similarity, the region of alignment, typically, exhibits at least about 75% of sequence identity; more 30 typically, at least about 78%; even more typically; at least about 80% sequence identity. Usually, percent sequence identity can be as much as about 82%; more usually, as much as about 84%; even more usually, as much as about 86%.

The p value is used in conjunction with these methods. If high similarity is found, the query sequence is considered to have high similarity with a profile 35

sequence when the p value is less than or equal to about 10⁻²; more usually; less than or equal to about 10⁻³; even more usually; less than or equal to about 10⁻⁴. More typically, the p value is no more than about 10⁻⁵; more typically; no more than or equal to about 10⁻¹⁰; even more typically; no more than or equal to about 10⁻¹⁵ for the query sequence to be considered high similarity.

Similarity Determined by Sequence Identity Alone. Sequence identity alone can be used to determine similarity of a query sequence to an individual sequence and can indicate the activity of the sequence. Such an alignment, preferably, permits gaps to align sequences. Typically, the query sequence is related to the profile sequence if the sequence identity over the entire query sequence is at least about 15%; more typically, at least about 20%; even more typically, at least about 50%. Sequence identity alone as a measure of similarity is most useful when the query sequence is usually, at least 80 residues in length; more usually, 90 residues; even more usually, at least 95 amino acid residues in length. More typically, similarity can be concluded based on sequence identity alone when the query sequence is preferably 100 residues in length; more preferably, 120 residues in length; even more preferably, 150 amino acid residues in length.

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Alignments with Profile and Multiple Aligned Sequences. Translations of the provided polynucleotides can be aligned with amino acid profiles that define either protein families or common motifs. Also, translations of the provided polynucleotides can be aligned to multiple sequence alignments (MSA) comprising the polypeptide sequences of members of protein families or motifs. Similarity or identity with profile sequences or MSAs can be used to determine the activity of the gene products (e.g., polypeptides) encoded by the provided polynucleotides or corresponding cDNA or genes. For example, sequences that show an identity or similarity with a chemokine profile or MSA can exhibit chemokine activities.

Profiles can be designed manually by (1) creating an MSA, which is an alignment of the amino acid sequence of members that belong to the family and (2) constructing a statistical representation of the alignment. Such methods are described, for example, in Birney et al., Nucl. Acid Res. (1996) 24(14): 2730-2739. MSAs of some protein families and motifs are publicly available. MSAs are described also in Sonnhammer et al., Proteins (1997) 28: 405-420. A brief description of MSAs is reported in Pascarella et al., Prot. Eng. (1996) 9(3):249-251. Techniques for building profiles from MSAs are described in Sonnhammer et al., supra; Birney et al., supra;

and "Computer Methods for Macromolecular Sequence Analysis," Methods in Enzymology (1996) 266, Doolittle, Academic Press, Inc., San Diego, California, USA.

Similarity between a query sequence and a protein family or motif can be determined by (a) comparing the query sequence against the profile and/or (b) aligning the query sequence with the members of the family or motif. Typically, a program such as Searchwise is used to compare the query sequence to the statistical representation of the multiple alignment, also known as a profile (see Birney et al., *supra*). Other techniques to compare the sequence and profile are described in Sonnhammer et al., *supra* and Doolittle, *supra*.

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Next, methods described by Feng et al., *J. Mol. Evol.* (1987) 25:351 and Higgins et al., *CABIOS* (1989) 5:151 can be used align the query sequence with the members of a family or motif, also known as a MSA. Sequence alignments can be generated using any of a variety of software tools. Examples include PileUp, which creates a multiple sequence alignment, and is described in Feng et al., *J. Mol. Evol.* (1987) 25:351. Another method, GAP, uses the alignment method of Needleman et al., *J. Mol. Biol.* (1970) 48:443. GAP is best suited for global alignment of sequences. A third method, BestFit, functions by inserting gaps to maximize the number of matches using the local homology algorithm of Smith et al., *Adv. Appl. Math.* (1981) 2:482. In general, the following factors are used to determine if a similarity between a query sequence and a profile or MSA exists: (1) number of conserved residues found in the query sequence, (2) percentage of conserved residues found in the query sequence, (3) number of frameshifts, and (4) spacing between conserved residues.

Some alignment programs that both translate and align sequences can make any number of frameshifts when translating the nucleotide sequence to produce the best alignment. The fewer frameshifts needed to produce an alignment, the stronger the similarity or identity between the query and profile or MSAs. For example, a weak similarity resulting from no frameshifts can be a better indication of activity or structure of a query sequence, than a strong similarity resulting from two frameshifts. Preferably, three or fewer frameshifts are found in an alignment; more preferably two or fewer frameshifts; even more preferably, one or fewer frameshifts; even more preferably, no frameshifts are found in an alignment of query and profile or MSAs.

Conserved residues are those amino acids found at a particular position in all or some of the family or motif members. Alternatively, a position is considered conserved if only a certain class of amino acids is found in a particular position in all or

some of the family members. For example, the N-terminal position can contain a positively charged amino acid, such as lysine, arginine, or histidine.

Typically, a residue of a polypeptide is conserved when a class of amino acids or a single amino acid is found at a particular position in at least about 40% of all class members; more typically, at least about 50%; even more typically, at least about 60% of the members. Usually, a residue is conserved when a class or single amino acid is found in at least about 70% of the members of a family or motif; more usually, at least about 80%; even more usually, at least about 90%; even more usually, at least about 95%.

A residue is considered conserved when three unrelated amino acids are found at a particular position in the some or all of the members; more usually, two unrelated amino acids. These residues are conserved when the unrelated amino acids are found at particular positions in at least about 40% of all class member; more typically, at least about 50%; even more typically, at least about 60% of the members. Usually, a residue is conserved when a class or single amino acid is found in at least about 70% of the members of a family or motif; more usually, at least about 80%; even more usually, at least about 90%; even more usually, at least about 95%.

A query sequence has similarity to a profile or MSA when the query sequence comprises at least about 25% of the conserved residues of the profile or MSA; more usually, at least about 30%; even more usually; at least about 40%. Typically, the query sequence has a stronger similarity to a profile sequence or MSA when the query sequence comprises at least about 45% of the conserved residues of the profile or MSA; more typically, at least about 50%; even more typically; at least about 55%.

Identification of Secreted and Membrane-Bound Polypeptides

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Both secreted and membrane-bound polypeptides of the present invention are of particular interest. For example, levels of secreted polypeptides can be assayed in body fluids that are convenient, such as blood, plasma, serum, and other body fluids such as urine, prostatic fluid and semen. Membrane-bound polypeptides are useful for constructing vaccine antigens or inducing an immune response. Such antigens would comprise all or part of the extracellular region of the membrane-bound polypeptides. Because both secreted and membrane-bound polypeptides comprise a fragment of contiguous hydrophobic amino acids, hydrophobicity predicting algorithms can be used to identify such polypeptides.

A signal sequence is usually encoded by both secreted and membrane-bound polypeptide genes to direct a polypeptide to the surface of the cell. The signal sequence usually comprises a stretch of hydrophobic residues. Such signal sequences can fold into helical structures. Membrane-bound polypeptides typically comprise at least one transmembrane region that possesses a stretch of hydrophobic amino acids that can transverse the membrane. Some transmembrane regions also exhibit a helical structure. Hydrophobic fragments within a polypeptide can be identified by using computer algorithms. Such algorithms include Hopp & Woods, *Proc. Natl. Acad. Sci. USA* (1981) 78:3824-3828; Kyte & Doolittle, *J. Mol. Biol.* (1982) 157: 105-132; and RAOAR algorithm, Degli Esposti et al., *Eur. J. Biochem.* (1990) 190: 207-219.

Another method of identifying secreted and membrane-bound polypeptides is to translate the polynucleotides of the invention in all six frames and determine if at least 8 contiguous hydrophobic amino acids are present. Those translated polypeptides with at least 8; more typically, 10; even more typically, 12 contiguous hydrophobic amino acids are considered to be either a putative secreted or membrane bound polypeptide. Hydrophobic amino acids include alanine, glycine, histidine, isolcucine, leucine, lysine, methionine, phenylalanine, proline, threonine, tryptophan, tyrosine, and valine

Identification of the Function of an Expression Product of a Full-Length Gene

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Ribozymes, antisense constructs, and dominant negative mutants can be used to determine function of the expression product of a gene corresponding to a polynucleotide provided herein. The phosphoramidite method of oligonucleotide synthesis can be used to construct antisense molecules and ribozymes. See Beaucage et al., *Tet. Lett.* (1981) 22:1859 and U.S. Patent No. 4,668,777. Automated devices for synthesis are available to create oligonucleotides using this chemistry. Examples of such devices include Biosearch 8600, Models 392 and 394 by Applied Biosystems, a division of Perkin-Elmer Corp., Foster City, California, USA; and Expedite by Perceptive Biosystems, Framingham, Massachusetts, USA. Synthetic RNA, phosphate analog oligonucleotides, and chemically derivatized oligonucleotides can also be produced, and can be covalently attached to other molecules. RNA oligonucleotides can be synthesized, for example, using RNA phosphoramidites. This method can be performed on an automated synthesizer, such as Applied Biosystems, Models 392 and 394, Foster City, California, USA.

Oligonucleotides of up to 200 nt can be synthesized, more typically, 100 nt, more typically 50 nt; even more typically 30 to 40 nt. These synthetic fragments can be annealed and ligated together to construct larger fragments. See, for example, Sambrook et al., *supra*. Trans-cleaving catalytic RNAs (ribozymes) are RNA molecules possessing endoribonuclease activity. Ribozymes are specifically designed for a particular target, and the target message must contain a specific nucleotide sequence. They are engineered to cleave any RNA species site-specifically in the background of cellular RNA. The cleavage event renders the mRNA unstable and prevents protein expression. Importantly, ribozymes can be used to inhibit expression of a gene of unknown function for the purpose of determining its function in an *in vitro* or *in vivo* context, by detecting the phenotypic effect.

Antisense nucleic acids are designed to specifically bind to RNA, resulting in the formation of RNA-DNA or RNA-RNA hybrids, with an arrest of DNA replication, reverse transcription or messenger RNA translation. Antisense polynucleotides based on a selected polynucleotide sequence can interfere with expression of the corresponding gene. Antisense polynucleotides are typically generated within the cell by expression from antisense constructs that contain the antisense strand as the transcribed strand. Antisense polynucleotides based on the disclosed polynucleotides will bind and/or interfere with the translation of mRNA comprising a sequence complementary to the antisense polynucleotide. The expression products of control cells and cells treated with the antisense construct are compared to detect the protein product of the gene corresponding to the polynucleotide upon which the antisense construct is based. The protein is isolated and identified using routine biochemical methods.

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Given the extensive background literature and clinical experience in antisense therapy, one skilled in the art can use selected polynucleotides of the invention as additional potential therapeutics. The choice of polynucleotide can be narrowed by first testing them for binding to "hot spot" regions of the genome of cancerous cells. If a polynucleotide is identified as binding to a "hot spot," testing the polynucleotide as an antisense compound in the corresponding cancer cells is warranted.

Dominant negative mutations also are readily generated for corresponding proteins that are active as homomultimers. A mutant polypeptide will interact with wild-type polypeptides (made from the other allele) and form a non-functional multimer. Thus, a mutation is in a substrate-binding domain, a catalytic

domain, or a cellular localization domain. Preferably, the mutant polypeptide will be overproduced. Point mutations are made that have such an effect. In addition, fusion of different polypeptides of various lengths to the terminus of a protein can yield dominant negative mutants. General strategies are available for making dominant negative mutants (see, e.g., Herskowitz, Nature (1987) 329:219). Such techniques can be used to create loss of function mutations, which are useful for determining protein function.

Polypeptides and Variants Thereof

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The polypeptides of the invention include those encoded by the disclosed polynucleotides, as well as nucleic acids that, by virtue of the degeneracy of the genetic code, are not identical in sequence to the disclosed polynucleotides. Thus, the invention includes within its scope a polypeptide encoded by a polynucleotide having the sequence of any one of SEQ ID NOs:1-3351 or a variant thereof.

In general, the term "polypeptide" as used herein refers to both the full length polypeptide encoded by the recited polynucleotide, the polypeptide encoded by the gene represented by the recited polynucleotide, as well as portions or fragments thereof. "Polypeptides" also includes variants of the naturally occurring proteins, where such variants are homologous or substantially similar to the naturally occurring protein, and can be of an origin of the same or different species as the naturally occurring protein (e.g., human, murine, or some other species that naturally expresses the recited polypeptide, usually a mammalian species). In general, variant polypeptides have a sequence that has at least about 80%, usually at least about 90%, and more usually at least about 98% sequence identity with a differentially expressed polypeptide of the invention, as measured by BLAST using the parameters described above. The variant polypeptides can be naturally or non-naturally glycosylated, i.e., the polypeptide has a glycosylation pattern that differs from the glycosylation pattern found in the corresponding naturally occurring protein.

The invention also encompasses homologs of the disclosed polypeptides (or fragments thereof) where the homologs are isolated from other species, i.e., other animal or plant species, where such homologs, usually mammalian species, e.g., rodents, such as mice, rats; domestic animals, e.g., horse, cow, dog, cat; and humans. By "homolog" is meant a polypeptide having at least about 35%, usually at least about 40% and more usually at least about 60% amino acid sequence identity to a particular differentially expressed protein as identified above, where sequence identity is determined using the BLAST algorithm, with the parameters described above.

In general, the polypeptides of the subject invention are provided in a non-naturally occurring environment, e.g., are separated from their naturally occurring environment. In certain embodiments, the subject protein is present in a composition that is enriched for the protein as compared to a control. As such, purified polypeptide is provided, where by purified is meant that the protein is present in a composition that is substantially free of non-differentially expressed polypeptides, where by substantially free is meant that less than 90%, usually less than 60% and more usually less than 50% of the composition is made up of non-differentially expressed polypeptides.

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Also within the scope of the invention are variants; variants of polypeptides include mutants, fragments, and fusions. Mutants can include amino acid substitutions, additions or deletions. The amino acid substitutions can be conservative amino acid substitutions or substitutions to eliminate non-essential amino acids, such as to alter a glycosylation site, a phosphorylation site or an acetylation site, or to minimize misfolding by substitution or deletion of one or more cysteine residues that are not necessary for function. Conservative amino acid substitutions are those that preserve the general charge, hydrophobicity/ hydrophilicity, and/or steric bulk of the amino acid substituted. Variants can be designed so as to retain biological activity of a particular region of the protein (e.g., a functional domain and/or, where the polypeptide is a member of a protein family, a region associated with a consensus sequence). Selection of amino acid alterations for production of variants can be based upon the accessibility (interior vs. exterior) of the amino acid (see, e.g., Go et al., Int. J. Peptide Protein Res. (1980) 15:211), the thermostability of the variant polypeptide (see, e.g., Querol et al., Prot. Eng. (1996) 9:265), desired glycosylation sites (see, e.g., Olsen and Thomsen, J. Gen. Microbiol. (1991) 137:579), desired disulfide bridges (see, e.g., Clarke et al., Biochemistry (1993) 32:4322; and Wakarchuk et al., Protein Eng. (1994) 7:1379), desired metal binding sites (see, e.g., Toma et al., Biochemistry (1991) 30:97, and Haezerbrouck et al., Protein Eng. (1993) 6:643), and desired substitutions with in proline loops (see, e.g., Masul et al., Appl. Env. Microbiol. (1994) 60:3579). Cysteinedepleted muteins can be produced as disclosed in U.S. Patent No. 4,959,314.

Variants also include fragments of the polypeptides disclosed herein, particularly biologically active fragments and/or fragments corresponding to functional domains. Fragments of interest will typically be at least about 10 aa to at least about 15 aa in length, usually at least about 50 aa in length, and can be as long as 300 aa in length or longer, but will usually not exceed about 1000 aa in length, where the fragment will have a stretch of amino acids that is identical to a polypeptide encoded by a

polynucleotide having a sequence of any SEQ ID NOs:1-3351, or a homolog thereof. The protein variants described herein are encoded by polynucleotides that are within the scope of the invention. The genetic code can be used to select the appropriate codons to construct the corresponding variants.

Computer-Related Embodiments

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In general, a library of polynucleotides is a collection of sequence information, which information is provided in either biochemical form (e.g., as a collection of polynucleotide molecules), or in electronic form (e.g., as a collection of polynucleotide sequences stored in a computer-readable form, as in a computer system The sequence information of the and/or as part of a computer program). polynucleotides can be used in a variety of ways, e.g., as a resource for genc discovery, as a representation of sequences expressed in a selected cell type (e.g., cell type markers), and/or as markers of a given disease or disease state. In general, a disease marker is a representation of a gene product that is present in all cells affected by disease either at an increased or decreased level relative to a normal cell (e.g., a cell of the same or similar type that is not substantially affected by disease). For example, a polynucleotide sequence in a library can be a polynucleotide that represents an mRNA, polypeptide, or other gene product encoded by the polynucleotide, that is either overexpressed or underexpressed in a breast ductal cell affected by cancer relative to a normal (i.e., substantially disease-free) breast cell. 20

The nucleotide sequence information of the library can be embodied in any suitable form, e.g., electronic or biochemical forms. For example, a library of sequence information embodied in electronic form comprises an accessible computer data file (or, in biochemical form, a collection of nucleic acid molecules) that contains the representative nucleotide sequences of genes that are differentially expressed (e.g., overexpressed or underexpressed) as between, for example, i) a cancerous cell and a normal cell; ii) a cancerous cell and a dysplastic cell; iii) a cancerous cell and a cell affected by a disease or condition other than cancer; iv) a metastatic cancerous cell and a normal cell and/or non-metastatic cancerous cell; v) a malignant cancerous cell and a non-malignant cancerous cell (or a normal cell) and/or vi) a dysplastic cell relative to a normal cell. Other combinations and comparisons of cells affected by various diseases or stages of disease will be readily apparent to the ordinarily skilled artisan. Biochemical embodiments of the library include a collection of nucleic acids that have

the sequences of the genes in the library, where the nucleic acids can correspond to the entire gene in the library or to a fragment thereof, as described in greater detail below.

The polynucleotide libraries of the subject invention generally comprise sequence information of a plurality of polynucleotide sequences, where at least one of the polynucleotides has a sequence of any of SEQ ID NOs:1-3351. By plurality is meant at least 2, usually at least 3 and can include up to all of SEQ ID NOs:1-3351. The length and number of polynucleotides in the library will vary with the nature of the library, e.g., if the library is an oligonucleotide array, a cDNA array, a computer database of the sequence information, etc.

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Where the library is an electronic library, the nucleic acid sequence information can be present in a variety of media. "Media" refers to a manufacture, other than an isolated nucleic acid molecule, that contains the sequence information of the present invention. Such a manufacture provides the genome sequence or a subset thereof in a form that can be examined by means not directly applicable to the sequence as it exists in a nucleic acid. For example, the nucleotide sequence of the present invention, e.g., the nucleic acid sequences of any of the polynucleotides of SEQ ID NOs:1-3351, can be recorded on computer readable media, e.g., any medium that can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as a floppy disc, a hard disc storage medium, and a magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. One of skill in the art can readily appreciate how any of the presently known computer readable mediums can be used to create a manufacture comprising a recording "Recorded" refers to a process for storing of the present sequence information. information on computer readable medium, using any such methods as known in the art. Any convenient data storage structure can be chosen, based on the means used to access the stored information. A variety of data processor programs and formats can be used for storage, e.g., word processing text file, database format, etc. In addition to the sequence information, electronic versions of the libraries of the invention can be provided in conjunction or connection with other computer-readable information and/or 30 other types of computer-readable files (e.g., searchable files, executable files, etc., including, but not limited to, for example, search program software, etc.).

By providing the nucleotide sequence in computer readable form, the information can be accessed for a variety of purposes. Computer software to access sequence information is publicly available. For example, the BLAST (Altschul et al.,

supra.) and BLAZE (Brutlag et al. Comp. Chem. (1993) 17:203) search algorithms on a Sybase system can be used to identify open reading frames (ORFs) within the genome that contain homology to ORFs from other organisms.

As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A skilled artisan can readily appreciate that any one of the currently available computer-based system are suitable for use in the present invention. The data storage means can comprise any manufacture comprising a recording of the present sequence information as described above, or a memory access means that can access such a manufacture.

"Search means" refers to one or more programs implemented on the computer-based system, to compare a target sequence or target structural motif, or expression levels of a polynucleotide in a sample, with the stored sequence information. Search means can be used to identify fragments or regions of the genome that match a particular target sequence or target motif. A variety of known algorithms are publicly known and commercially available, e.g., MacPattern (EMBL), BLASTN and BLASTX (NCBI). A "target sequence" can be any polynucleotide or amino acid sequence of six or more contiguous nucleotides or two or more amino acids, preferably from about 10 to 100 amino acids or from about 30 to 300 nt. A variety of comparing means can be used to accomplish comparison of sequence information from a sample (e.g., to analyze target sequences, target motifs, or relative expression levels) with the data storage means. A skilled artisan can readily recognize that any one of the publicly available homology search programs can be used as the search means for the computer based systems of the present invention to accomplish comparison of target sequences and motifs. Computer programs to analyze expression levels in a sample and in controls are also known in the art.

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A "target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) are chosen based on a three-dimensional configuration that is formed upon the folding of the target motif, or on consensus sequences of regulatory or active sites. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzyme active sites and signal sequences. Nucleic acid target motifs include, but are

WO 01/02568

not limited to, hairpin structures, promoter sequences and other expression elements such as binding sites for transcription factors.

A variety of structural formats for the input and output means can be used to input and output the information in the computer-based systems of the present invention. One format for an output means ranks the relative expression levels of different polynucleotides. Such presentation provides a skilled artisan with a ranking of relative expression levels to determine a gene expression profile.

As discussed above, the "library" of the invention also encompasses biochemical libraries of the polynucleotides of SEQ ID NOs:1-3351, e.g., collections of nucleic acids representing the provided polynucleotides. The biochemical libraries can take a variety of forms, e.g., a solution of cDNAs, a pattern of probe nucleic acids stably associated with a surface of a solid support (i.e., an array) and the like. Of particular interest are nucleic acid arrays in which one or more of SEQ ID NOs:1-3351 is represented on the array. By array is meant an article of manufacture that has at least a substrate with at least two distinct nucleic acid targets on one of its surfaces, where the number of distinct nucleic acids can be considerably higher, typically being at least 10 nt, usually at least 20 nt and often at least 25 nt. A variety of different array formats have been developed and are known to those of skill in the art. The arrays of the subject invention find use in a variety of applications, including gene expression analysis, drug screening, mutation analysis and the like, as disclosed in the above-listed exemplary patent documents.

In addition to the above nucleic acid libraries, analogous libraries of polypeptides are also provided, where the where the polypeptides of the library will represent at least a portion of the polypeptides encoded by SEQ ID NOs:1-3351.

25 Use of Polynucleotide Probes in Mapping, and in Tissue Profiling

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Polynucleotide probes, generally comprising at least 12 contiguous nt of a polynucleotide as shown in the Sequence Listing, are used for a variety of purposes, such as chromosome mapping of the polynucleotide and detection of transcription levels. Additional disclosure about preferred regions of the disclosed polynucleotide sequences is found in the Examples. A probe that hybridizes specifically to a polynucleotide disclosed herein should provide a detection signal at least 5-, 10-, or 20-fold higher than the background hybridization provided with other unrelated sequences.

<u>Detection of Expression Levels</u>. Nucleotide probes are used to detect expression of a gene corresponding to the provided polynucleotide. In Northern blots,

mRNA is separated electrophoretically and contacted with a probe. A probe is detected as hybridizing to an mRNA species of a particular size. The amount of hybridization is quantitated to determine relative amounts of expression, for example under a particular condition. Probes are used for *in situ* hybridization to cells to detect expression. Probes can also be used *in vivo* for diagnostic detection of hybridizing sequences. Probes are typically labeled with a radioactive isotope. Other types of detectable labels can be used such as chromophores, fluors, and enzymes. Other examples of nucleotide hybridization assays are described in WO92/02526 and U.S. Patent No. 5,124,246.

Alternatively, the Polymerase Chain Reaction (PCR) is another means for detecting small amounts of target nucleic acids (see, e.g., Mullis et al., Meth. Enzymol. (1987) 155:335; U.S. Patent No. 4,683,195; and U.S. Patent No. 4,683,202). Two primer polynucleotides nucleotides that hybridize with the target nucleic acids are used to prime the reaction. The primers can be composed of sequence within or 3' and 5' to the polynucleotides of the Sequence Listing. Alternatively, if the primers are 3' and 5' to these polynucleotides, they need not hybridize to them or the complements. After 15 amplification of the target with a thermostable polymerase, the amplified target nucleic acids can be detected by methods known in the art, e.g., Southern blot. mRNA or cDNA can also be detected by traditional blotting techniques (e.g., Southern blot, Northern blot, etc.) described in Sambrook et al., "Molecular Cloning: A Laboratory Manual" (New York, Cold Spring Harbor Laboratory, 1989) (e.g., without PCR 20 amplification). In general, mRNA or cDNA generated from mRNA using a polymerase enzyme can be purified and separated using gel electrophoresis, and transferred to a solid support, such as nitrocellulose. The solid support is exposed to a labeled probe, washed to remove any unhybridized probe, and duplexes containing the labeled probe are detected. 25

Mapping. Polynucleotides of the present invention can be used to identify a chromosome on which the corresponding gene resides. Such mapping can be useful in identifying the function of the polynucleotide-related gene by its proximity to other genes with known function. Function can also be assigned to the polynucleotide-related gene when particular syndromes or diseases map to the same chromosome. For example, use of polynucleotide probes in identification and quantification of nucleic acid sequence aberrations is described in U.S. Patent No. 5,783,387. An exemplary mapping method is fluorescence in situ hybridization (FISH), which facilitates comparative genomic hybridization to allow total genome assessment of changes in relative copy number of DNA sequences (see, e.g., Valdes et al., Methods in Molecular

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Biology (1997) 68:1). Polynucleotides can also be mapped to particular chromosomes using, for example, radiation hybrids or chromosome-specific hybrid panels. See Leach et al., Advances in Genetics, (1995) 33:63-99; Walter et al., Nature Genetics (1994) 7:22; Walter and Goodfellow, Trends in Genetics (1992) 9:352. Panels for radiation hybrid mapping are available from Research Genetics, Inc., Huntsville, Alabama, USA. The statistical program RHMAP can be used to construct a map based on the data from radiation hybridization with a measure of the relative likelihood of one order versus another. RHMAP is available via the world wide web at http://www.sph.umich.edu/group/statgen/software. In addition, commercial programs are available for identifying regions of chromosomes commonly associated with disease, such as cancer.

Tissue Typing or Profiling. Expression of specific mRNA corresponding to the provided polynucleotides can vary in different cell types and can be tissue-specific. This variation of mRNA levels in different cell types can be exploited with nucleic acid probe assays to determine tissue types. For example, PCR, branched DNA probe assays, or blotting techniques utilizing nucleic acid probes substantially identical or complementary to polynucleotides listed in the Sequence Listing can determine the presence or absence of the corresponding cDNA or mRNA.

Tissue typing can be used to identify the developmental organ or tissue source of a metastatic lesion by identifying the expression of a particular marker of that organ or tissue. If a polynucleotide is expressed only in a specific tissue type, and a metastatic lesion is found to express that polynucleotide, then the developmental source of the lesion has been identified. Expression of a particular polynucleotide can be assayed by detection of either the corresponding mRNA or the protein product.

<u>Use of Polymorphisms</u>. A polynucleotide of the invention can be used in forensics, genetic analysis, mapping, and diagnostic applications where the corresponding region of a gene is polymorphic in the human population. Any means for detecting a polymorphism in a gene can be used, including, but not limited to electrophoresis of protein polymorphic variants, differential sensitivity to restriction enzyme cleavage, and hybridization to allele-specific probes.

30 Antibody Production

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Expression products of a polynucleotide of the invention, as well as the corresponding mRNA, cDNA, or complete gene, can be prepared and used for raising antibodies for experimental, diagnostic, and therapeutic purposes. For polynucleotides to which a corresponding gene has not been assigned, this provides an additional

method of identifying the corresponding gene. The polynucleotide or related cDNA is expressed as described above, and antibodies are prepared. These antibodies are specific to an epitope on the polypeptide encoded by the polynucleotide, and can precipitate or bind to the corresponding native protein in a cell or tissue preparation or in a cell-free extract of an *in vitro* expression system.

Methods for production of monoclonal and polyclonal antibodies that specifically bind a selected antigen are well known in the art. The antibodies specifically bind to epitopes present in the polypeptides encoded by polynucleotides disclosed in the Sequence Listing. Typically, at least 6, 8, 10, or 12 contiguous amino acids are required to form an cpitope. Epitopes that involve non-contiguous amino acids may require a longer polypeptide, e.g., at least 15, 25, or 50 amino acids. Antibodies that specifically bind to human polypeptides encoded by the provided polynucleotides should provide a detection signal at least 5-, 10-, or 20-fold higher than a detection signal provided with other proteins when used in Western blots or other immunochemical assays. Preferably, antibodies that specifically polypeptides of the invention do not bind to other proteins in immunochemical assays at detectable levels and can immunoprecipitate the specific polypeptide from solution.

The invention also contemplates naturally occurring antibodies specific for a polypeptide of the invention. For example, serum antibodies to a polypeptide of the invention in a human population can be purified by methods well known in the art, e.g., by passing antiserum over a column to which the corresponding selected polypeptide or fusion protein is bound. The bound antibodies can then be eluted from the column, for example using a buffer with a high salt concentration.

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In addition to the antibodies discussed above, the invention also contemplates genetically engineered antibodies, antibody derivatives (e.g., single chain antibodies, antibody fragments (e.g., Fab, etc.)), according to methods well known in the art.

Other embodiments of the present invention include humanized monoclonal antibodies capable of binding to the polypeptides of the invention. The phrase "humanized antibody" refers to an antibody derived from a non-human antibody - typically a mouse monoclonal antibody. Alternatively, a humanized antibody may be derived from a chimeric antibody that retains or substantially retains the antigen-binding properties of the parental, non-human, antibody but which exhibits diminished immunogenicity as compared to the parental antibody when administered to humans. The phrase "chimeric antibody," as used herein, refers to an antibody containing

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sequence derived from two different antibodies (see, e.g., U.S. Patent No. 4,816,567) which typically originate from different species. Most typically, chimeric antibodies comprise human and murine antibody fragments, generally human constant and mouse variable regions.

Because humanized antibodies are far less immunogenic in humans than the parental mouse monoclonal antibodies, they can be used for the treatment of humans with far less risk of anaphylaxis. Thus, these antibodies may be preferred in therapeutic applications that involve *in vivo* administration to a human such as, *e.g.*, use as radiation sensitizers for the treatment of neoplastic disease or use in methods to reduce the side effects of, *e.g.*, cancer therapy.

Humanized antibodies may be achieved by a variety of methods including, for example: (1) grafting the non-human complementarity determining regions (CDRs) onto a human framework and constant region (a process referred to in the art as "humanizing"), or, alternatively, (2) transplanting the entire non-human variable domains, but "cloaking" them with a human-like surface by replacement of surface residues (a process referred to in the art as "veneering"). In the present invention, humanized antibodies will include both "humanized" and "veneered" antibodies. These methods are disclosed in, e.g., Jones et al., Nature 321:522-525 (1986); Morrison et al., Proc. Natl. Acad. Sci., U.S.A., 81:6851-6855 (1984); Morrison and Oi, Adv. Immunol., 44:65-92 (1988); Verhoeyer et al., Science 239:1534-1536 (1988); Padlan, Molec. Immun. 28:489-498 (1991); Padlan, Molec. Immunol. 31(3):169-217 (1994); and Kettleborough, C.A. et al., Protein Eng. 4(7):773-83 (1991) each of which is incorporated herein by reference.

The phrase "complementarity determining region" refers to amino acid sequences which together define the binding affinity and specificity of the natural Fv region of a native immunoglobulin binding site. See, e.g., Chothia et al., J. Mol. Biol. 196:901-917 (1987); Kabat et al., U.S. Dept. of Health and Human Services NIH Publication No. 91-3242 (1991). The phrase "constant region" refers to the portion of the antibody molecule that confers effector functions. In the present invention, mouse constant regions are substituted by human constant regions. The constant regions of the subject humanized antibodies are derived from human immunoglobulins. The heavy chain constant region can be selected from any of the five isotypes: alpha, delta, epsilon, gamma or mu.

One method of humanizing antibodies comprises aligning the nonhuman heavy and light chain sequences to human heavy and light chain sequences,

selecting and replacing the non-human framework with a human framework based on such alignment, molecular modeling to predict the conformation of the humanized sequence and comparing to the conformation of the parent antibody. This process is followed by repeated back mutation of residues in the CDR region which disturb the structure of the CDRs until the predicted conformation of the humanized sequence model closely approximates the conformation of the non-human CDRs of the parent non-human antibody. Such humanized antibodies may be further derivatized to facilitate uptake and clearance, e.g., via Ashwell receptors. See, e.g., U.S. Patent Nos. 5,530,101 and 5,585,089 which patents are incorporated herein by reference.

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Humanized antibodies can also be produced using transgenic animals that are engineered to contain human immunoglobulin loci. For example, WO 98/24893 discloses transgenic animals having a human Ig locus wherein the animals do not produce functional endogenous immunoglobulins due to the inactivation of endogenous heavy and light chain loci. WO 91/10741 also discloses transgenic non-primate mammalian hosts capable of mounting an immune response to an immunogen, wherein the antibodies have primate constant and/or variable regions, and wherein the endogenous immunoglobulin-encoding loci are substituted or inactivated. WO 96/30498 discloses the use of the Cre/Lox system to modify the immunoglobulin locus in a mammal, such as to replace all or a portion of the constant or variable region to form a modified antibody molecule. WO 94/02602 discloses non-human mammalian hosts having inactivated endogenous Ig loci and functional human Ig loci. U.S. Patent No. 5,939,598 discloses methods of making transgenic mice in which the mice lack endogenous heavy claims, and express an exogenous immunoglobulin locus comprising one or more xenogeneic constant regions.

Using a transgenic animal described above, an immune response can be produced to a selected antigenic molecule, and antibody-producing cells can be removed from the animal and used to produce hybridomas that secrete human monoclonal antibodies. Immunization protocols, adjuvants, and the like are known in the art, and are used in immunization of, for example, a transgenic mouse as described in WO 96/33735. This publication discloses monoclonal antibodies against a variety of antigenic molecules including IL-6, IL-8, TNF, human CD4, L-selectin, gp39, and tetanus toxin. The monoclonal antibodies can be tested for the ability to inhibit or neutralize the biological activity or physiological effect of the corresponding protein. WO 96/33735 discloses that monoclonal antibodies against IL-8, derived from immune cells of transgenic mice immunized with IL-8, blocked IL-8-induced functions of

Human monoclonal antibodies with specificity for the antigen used to neutrophils. immunize transgenic animals are also disclosed in WO 96/34096.

Polynucleotides or Arrays for Diagnostics

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Polynucleotide arrays are created by spotting polynucleotide probes onto a substrate (e.g., glass, nitrocellose, etc.) in a two-dimensional matrix or array having bound probes. The probes can be bound to the substrate by either covalent bonds or by non-specific interactions, such as hydrophobic interactions. Samples of polynucleotides can be detectably labeled (e.g., using radioactive or fluorescent labels) and then hybridized to the probes. Double stranded polynucleotides, comprising the labeled sample polynucleotides bound to probe polynucleotides, can be detected once the unbound portion of the sample is washed away. Techniques for constructing arrays and methods of using these arrays are described in EP 799 897; WO 97/29212; WO 97/27317; EP 785 280; WO 97/02357; U.S. Patent No. 5,593,839; U.S. Patent No. 5,578,832; EP 728 520; U.S. Patent No. 5,599,695; EP 721 016; U.S. Patent No. 5,556,752; WO 95/22058; and U.S. Patent No. 5,631,734. Arrays can be used to, for example, examine differential expression of genes and can be used to determine gene function. For example, arrays can be used to detect differential expression of a polynucleotide between a test cell and control cell (e.g., cancer cells and normal cells). For example, high expression of a particular message in a cancer cell, which is not 20 observed in a corresponding normal cell, can indicate a cancer specific gene product. Exemplary uses of arrays are further described in, for example, Pappalarado et al., Sem. Radiation Oncol. (1998) 8:217; and Ramsay, Nature Biotechnol. (1998) 16:40.

Differential Expression in Diagnosis

The polynucleotides of the invention can also be used to detect differences in expression levels between two cells, e.g., as a method to identify abnormal or diseased tissue in a human. For polynucleotides corresponding to profiles of protein families, the choice of tissue can be selected according to the putative biological function. In general, the expression of a gene corresponding to a specific polynucleotide is compared between a first tissue that is suspected of being diseased and a second, normal tissue of the human. The tissue suspected of being abnormal or diseased can be derived from a different tissue type of the human, but preferably it is derived from the same tissue type; for example an intestinal polyp or other abnormal growth should be compared with normal intestinal tissue. The normal tissue can be the

same tissue as that of the test sample, or any normal tissue of the patient, especially those that express the polynucleotide-related gene of interest (e.g., brain, thymus, testis, heart, prostate, placenta, spleen, small intestine, skeletal muscle, pancreas, and the mucosal lining of the colon). A difference between the polynucleotide-related gene, mRNA, or protein in the two tissues which are compared, for example in molecular weight, amino acid or nucleotide sequence, or relative abundance, indicates a change in the gene, or a gene which regulates it, in the tissue of the human that was suspected of being diseased. Examples of detection of differential expression and its use in diagnosis of cancer are described in U.S. Patent Nos. 5,688,641 and 5,677,125.

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A genetic predisposition to disease in a human can also be detected by comparing expression levels of an mRNA or protein corresponding to a polynucleotide of the invention in a fetal tissue with levels associated in normal fetal tissue. Fetal tissues that are used for this purpose include, but are not limited to, amniotic fluid, chorionic villi, blood, and the blastomere of an in vitro-fertilized embryo. comparable normal polynucleotide-related gene is obtained from any tissue. The mRNA or protein is obtained from a normal tissue of a human in which the polynucleotiderelated gene is expressed. Differences such as alterations in the nucleotide sequence or size of the same product of the fétal polynucleotide-related gene or mRNA, or alterations in the molecular weight, amino acid sequence, or relative abundance of fetal protein, can indicate a germline mutation in the polynucleotide-related gene of the fetus, which indicates a genetic predisposition to disease. In general, diagnostic, prognostic, and other methods of the invention based on differential expression involve detection of a level or amount of a gene product, particularly a differentially expressed gene product, in a test sample obtained from a patient suspected of having or being susceptible to a disease (e.g., breast cancer, lung cancer, colon cancer and/or metastatic forms thereof), and comparing the detected levels to those levels found in normal cells (e.g., cells substantially unaffected by cancer) and/or other control cells (e.g., to differentiate a cancerous cell from a cell affected by dysplasia). Furthermore, the severity of the disease can be assessed by comparing the detected levels of a differentially expressed gene product with those levels detected in samples representing the levels of 30 differentially gene product associated with varying degrees of severity of disease. It should be noted that use of the term "diagnostic" herein is not necessarily meant to exclude "prognostic" or "prognosis," but rather is used as a matter of convenience.

The term "differentially expressed gene" is generally intended to encompass a polynucleotide that can, for example, include an open reading frame

encoding a gene product (e.g., a polypeptide), and/or introns of such genes and adjacent 5' and 3' non-coding nucleotide sequences involved in the regulation of expression, up to about 20 kb beyond the coding region, but possibly further in either direction. The gene can be introduced into an appropriate vector for extrachromosomal maintenance or for integration into a host genome. In general, a difference in expression level associated with a decrease in expression level of at least about 25%, usually at least about 50% to 75%, more usually at least about 90% or more is indicative of a differentially expressed gene of interest, i.e., a gene that is underexpressed or down-regulated in the test sample relative to a control sample. Furthermore, a difference in expression level associated with an increase in expression of at least about 25%, usually at least about 50% to 75%, more usually at least about 90% and can be at least about 1½-fold, usually at least about 2-fold to about 10-fold, and can be about 100-fold to about 1,000-fold increase relative to a control sample is indicative of a differentially expressed gene of interest, i.e., an overexpressed or up-regulated gene.

"Differentially expressed polynucleotide" as used herein means a nucleic acid molecule (RNA or DNA) comprising a sequence that represents a differentially expressed gene, e.g., the differentially expressed polynucleotide comprises a sequence (e.g., an open reading frame encoding a gene product) that uniquely identifies a differentially expressed gene so that detection of the differentially expressed polynucleotide in a sample is correlated with the presence of a differentially expressed gene in a sample. "Differentially expressed polynucleotides" is also meant to encompass fragments of the disclosed polynucleotides, e.g., fragments retaining biological activity, as well as nucleic acids homologous, substantially similar, or substantially identical (e.g., having about 90% sequence identity) to the disclosed polynucleotides.

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"Diagnosis" as used herein generally includes determination of a subject's susceptibility to a disease or disorder, determination as to whether a subject is presently affected by a disease or disorder, as well as to the prognosis of a subject affected by a disease or disorder (e.g., identification of pre-metastatic or metastatic cancerous states, stages of cancer, or responsiveness of cancer to therapy). The present invention particularly encompasses diagnosis of subjects in the context of breast cancer (e.g., carcinoma in situ (e.g., ductal carcinoma in situ), estrogen receptor (ER)-positive breast cancer, ER-negative breast cancer, or other forms and/or stages of breast cancer), lung cancer (e.g., small cell carcinoma, non-small cell carcinoma, mesothelioma, and

other forms and/or stages of lung cancer), and colon cancer (e.g., adenomatous polyp, colorectal carcinoma, and other forms and/or stages of colon cancer).

"Sample" or "biological sample" as used throughout here are generally meant to refer to samples of biological fluids or tissues, particularly samples obtained from tissues, especially from cells of the type associated with the disease for which the diagnostic application is designed (e.g., ductal adenocarcinoma), and the like. "Samples" is also meant to encompass derivatives and fractions of such samples (e.g., cell lysates). Where the sample is solid tissue, the cells of the tissue can be dissociated or tissue sections can be analyzed.

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Methods of the subject invention useful in diagnosis or prognosis typically involve comparison of the abundance of a selected differentially expressed gene product in a sample of interest with that of a control to determine any relative differences in the expression of the gene product, where the difference can be measured qualitatively and/or quantitatively. Quantitation can be accomplished, for example, by comparing the level of expression product detected in the sample with the amounts of product present in a standard curve. A comparison can be made visually; by using a technique such as densitometry, with or without computerized assistance; by preparing a representative library of cDNA clones of mRNA isolated from a test sample, sequencing the clones in the library to determine that number of cDNA clones corresponding to the same gene product, and analyzing the number of clones corresponding to that same gene product relative to the number of clones of the same gene product in a control sample; or by using an array to detect relative levels of hybridization to a selected sequence or set of sequences, and comparing the hybridization pattern to that of a control. The differences in expression are then correlated with the presence or absence of an abnormal expression pattern. A variety of different methods for determining the nucleic acid abundance in a sample are known to those of skill in the art (see, e.g., WO 97/27317). In general, diagnostic assays of the invention involve detection of a gene product of a the polynucleotide sequence (e.g., mRNA or polypeptide) that corresponds to a sequence of SEQ ID NOs:1-3351. The patient from whom the sample is obtained can be apparently healthy, susceptible to disease (e.g., as determined by family history or exposure to certain environmental factors), or can already be identified as having a condition in which altered expression of a gene product of the invention is implicated.

Diagnosis can be determined based on detected gene product expression levels of a gene product encoded by at least one, preferably at least two or more, at least

3 or more, or at least 4 or more of the polynucleotides having a sequence set forth in SEQ ID NOs:1-3351, and can involve detection of expression of genes corresponding to all of SEQ ID NOs:1-3351 and/or additional sequences that can serve as additional diagnostic markers and/or reference sequences. Where the diagnostic method is designed to detect the presence or susceptibility of a patient to cancer, the assay preferably involves detection of a gene product encoded by a gene corresponding to a polynucleotide that is differentially expressed in cancer. Examples of such differentially expressed polynucleotides are described in the Examples below. Given the provided polynucleotides and information regarding their relative expression levels provided herein, assays using such polynucleotides and detection of their expression levels in diagnosis and prognosis will be readily apparent to the ordinarily skilled artisan.

Any of a variety of detectable labels can be used in connection with the various embodiments of the diagnostic methods of the invention. Suitable detectable labels include fluorochromes, (e.g., fluorescein isothiocyanate (FITC), rhodamine, Texas Red, phycoerythrin, allophycocyanin, 6-carboxyfluorescein (6-FAM), 2',7'-dimethoxy-4',5'-dichloro-6-carboxyfluorescein, 6-carboxy-X-rhodamine (ROX), 6-carboxy-2',4',7',4,7-hexachlorofluorescein (HEX), 5-carboxyfluorescein (5-FAM) or N,N,N',N'-tetramethyl-6-carboxyrhodamine (TAMRA)), radioactive labels, (e.g., ³²P, ³⁵S, ³H, etc.), and the like. The detectable label can involve a two stage systems (e.g., biotin-avidin, hapten-anti-hapten antibody, etc.)

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Reagents specific for the polynucleotides and polypeptides of the invention, such as antibodies and nucleotide probes, can be supplied in a kit for detecting the presence of an expression product in a biological sample. The kit can also contain buffers or labeling components, as well as instructions for using the reagents to detect and quantify expression products in the biological sample. Exemplary embodiments of the diagnostic methods of the invention are described below in more detail.

Polypeptide detection in diagnosis. In one embodiment, the test sample is assayed for the level of a differentially expressed polypeptide. Diagnosis can be accomplished using any of a number of methods to determine the absence or presence or altered amounts of the differentially expressed polypeptide in the test sample. For example, detection can utilize staining of cells or histological sections with labeled antibodies, performed in accordance with conventional methods. Cells can be permeabilized to stain cytoplasmic molecules. In general, antibodies that specifically bind a differentially expressed polypeptide of the invention are added to a sample, and

incubated for a period of time sufficient to allow binding to the epitope, usually at least about 10 minutes. The antibody can be detectably labeled for direct detection (e.g., using radioisotopes, enzymes, fluorescers, chemiluminescers, and the like), or can be used in conjunction with a second stage antibody or reagent to detect binding (e.g., biotin with horseradish peroxidase-conjugated avidin, a secondary antibody conjugated to a fluorescent compound, e.g., fluorescein, rhodamine, Texas red, etc.). The absence or presence of antibody binding can be determined by various methods, including flow cytometry of dissociated cells, microscopy, radiography, scintillation counting, etc. Any suitable alternative methods can of qualitative or quantitative detection of levels or amounts of differentially expressed polypeptide can be used, for example ELISA, western blot, immunoprecipitation, radioimmunoassay, etc.

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mRNA detection. The diagnostic methods of the invention can also or alternatively involve detection of mRNA encoded by a gene corresponding to a differentially expressed polynucleotides of the invention. Any suitable qualitative or quantitative methods known in the art for detecting specific mRNAs can be used. mRNA can be detected by, for example, in situ hybridization in tissue sections, by reverse transcriptase-PCR, or in Northern blots containing poly A+ mRNA. One of skill in the art can readily use these methods to determine differences in the size or amount of mRNA transcripts between two samples. mRNA expression levels in a sample can also be determined by generation of a library of expressed scquence tags (ESTs) from the sample, where the EST library is representative of sequences present in the sample (Adams, et al., (1991) Science 252:1651). Enumeration of the relative representation of ESTs within the library can be used to approximate the relative representation of the gene transcript within the starting sample. The results of EST analysis of a test sample can then be compared to EST analysis of a reference sample to determine the relative expression levels of a selected polynucleotide, particularly a polynucleotide corresponding to one or more of the differentially expressed genes described herein. Alternatively, gene expression in a test sample can be performed using serial analysis of gene expression (SAGE) methodology (e.g., Velculescu et al., Science (1995) 270:484) or differential display (DD) methodology (see, e.g., U.S. Patent NOs. 5,776,683 and 5,807,680).

Alternatively, gene expression can be analyzed using hybridization analysis. Oligonucleotides or cDNA can be used to selectively identify or capture DNA or RNA of specific sequence composition, and the amount of RNA or cDNA hybridized to a known capture sequence determined qualitatively or quantitatively, to provide

information about the relative representation of a particular message within the pool of cellular messages in a sample. Hybridization analysis can be designed to allow for concurrent screening of the relative expression of hundreds to thousands of genes by using, for example, array-based technologies having high density formats, including filters, microscope slides, or microchips, or solution-based technologies that use spectroscopic analysis (e.g., mass spectrometry). One exemplary use of arrays in the diagnostic methods of the invention is described below in more detail.

Use of a single gene in diagnostic applications. The diagnostic methods of the invention can focus on the expression of a single differentially expressed gene. For example, the diagnostic method can involve detecting a differentially expressed gene, or a polymorphism of such a gene (e.g., a polymorphism in an coding region or control region), that is associated with disease. Disease-associated polymorphisms can include deletion or truncation of the gene, mutations that alter expression level and/or affect activity of the encoded protein, etc.

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A number of methods are available for analyzing nucleic acids for the presence of a specific sequence, e.g., a disease associated polymorphism. Where large amounts of DNA are available, genomic DNA is used directly. Alternatively, the region of interest is cloned into a suitable vector and grown in sufficient quantity for analysis. Cells that express a differentially expressed gene can be used as a source of mRNA, which can be assayed directly or reverse transcribed into cDNA for analysis. The nucleic acid can be amplified by conventional techniques, such as the polymerase chain reaction (PCR), to provide sufficient amounts for analysis, and a detectable label can be included in the amplification reaction (e.g., using a detectably labeled primer or detectably labeled oligonucleotides) to facilitate detection. Alternatively, various methods are also known in the art that utilize oligonucleotide ligation as a means of detecting polymorphisms, see e.g., Riley et al., Nucl. Acids Res. (1990) 18:2887; and Delahunty et al., Am. J. Hum. Genet. (1996) 58:1239.

The amplified or cloned sample nucleic acid can be analyzed by one of a number of methods known in the art. The nucleic acid can be sequenced by dideoxy or other methods, and the sequence of bases compared to a selected sequence, e.g., to a wild-type sequence. Hybridization with the polymorphic or variant sequence can also be used to determine its presence in a sample (e.g., by Southern blot, dot blot, etc.). The hybridization pattern of a polymorphic or variant sequence and a control sequence to an array of oligonucleotide probes immobilized on a solid support, as described in U.S. Patent No. 5,445,934, or in WO 95/35505, can also be used as a means of identifying

polymorphic or variant sequences associated with disease. Single strand conformational polymorphism (SSCP) analysis, denaturing gradient gel electrophoresis (DGGE), and heteroduplex analysis in gel matrices are used to detect conformational changes created by DNA sequence variation as alterations in electrophoretic mobility. Alternatively, where a polymorphism creates or destroys a recognition site for a restriction endonuclease, the sample is digested with that endonuclease, and the products size fractionated to determine whether the fragment was digested. Fractionation is performed by gel or capillary electrophoresis, particularly acrylamide or agarose gels.

Screening for mutations in a gene can be based on the functional or antigenic characteristics of the protein. Protein truncation assays are useful in detecting deletions that can affect the biological activity of the protein. Various immunoassays designed to detect polymorphisms in proteins can be used in screening. Where many diverse genetic mutations lead to a particular disease phenotype, functional protein assays have proven to be effective screening tools. The activity of the encoded protein can be determined by comparison with the wild-type protein.

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Pattern matching in diagnosis using arrays. In another embodiment, the diagnostic and/or prognostic methods of the invention involve detection of expression of a selected set of genes in a test sample to produce a test expression pattern (TEP). The TEP is compared to a reference expression pattern (REP), which is generated by detection of expression of the selected set of genes in a reference sample (e.g., a positive or negative control sample). The selected set of genes includes at least one of the genes of the invention, which genes correspond to the polynucleotide sequences of SEQ ID NOs:1-3351. Of particular interest is a selected set of genes that includes genes differentially expressed in the disease for which the test sample is to be screened.

"Reference sequences" or "reference polynucleotides" as used herein in the context of differential gene expression analysis and diagnosis/prognosis refers to a selected set of polynucleotides, which selected set includes at least one or more of the differentially expressed polynucleotides described herein. A plurality of reference sequences, preferably comprising positive and negative control sequences, can be included as reference sequences. Additional suitable reference sequences are found in Genbank, Unigene, and other nucleotide sequence databases (including, e.g., expressed sequence tag (EST), partial, and full-length sequences).

"Reference array" means an array having reference sequences for use in hybridization with a sample, where the reference sequences include all, at least one of,

or any subset of the differentially expressed polynucleotides described herein. Usually such an array will include at least 3 different reference sequences, and can include any one or all of the provided differentially expressed sequences. Arrays of interest can further comprise sequences, including polymorphisms, of other genetic sequences, particularly other sequences of interest for screening for a disease or disorder (e.g., cancer, dysplasia, or other related or unrelated diseases, disorders, or conditions). The oligonucleotide sequence on the array will usually be at least about 12 nt in length, and can be of about the length of the provided sequences, or can extend into the flanking regions to generate fragments of 100 nt to 200 nt in length or more. Reference arrays can be produced according to any suitable methods known in the art. For example, methods of producing large arrays of oligonucleotides are described in U.S. Patent NOs. 5,134,854 and 5,445,934 using light-directed synthesis techniques. Using a computer controlled system, a heterogeneous array of monomers is converted, through simultaneous coupling at a number of reaction sites, into a heterogeneous array of polymers. Alternatively, microarrays are generated by deposition of pre-synthesized oligonucleotides onto a solid substrate, for example as described in PCT published application no. WO 95/35505.

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A "reference expression pattern" or "REP" as used herein refers to the relative levels of expression of a selected set of genes, particularly of differentially expressed genes, that is associated with a selected cell type, e.g., a normal cell, a cancerous cell, a cell exposed to an environmental stimulus, and the like. A "test expression pattern" or "TEP" refers to relative levels of expression of a selected set of genes, particularly of differentially expressed genes, in a test sample (e.g., a cell of unknown or suspected disease state, from which mRNA is isolated).

REPs can be generated in a variety of ways according to methods well known in the art. For example, REPs can be generated by hybridizing a control sample to an array having a selected set of polynucleotides (particularly a selected set of differentially expressed polynucleotides), acquiring the hybridization data from the array, and storing the data in a format that allows for ready comparison of the REP with a TEP. Alternatively, all expressed sequences in a control sample can be isolated and sequenced, e.g., by isolating mRNA from a control sample, converting the mRNA into cDNA, and sequencing the cDNA. The resulting sequence information roughly or precisely reflects the identity and relative number of expressed sequences in the sample. The sequence information can then be stored in a format (e.g., a computer-readable format) that allows for ready comparison of the REP with a TEP. The REP can be

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normalized prior to or after data storage, and/or can be processed to selectively remove sequences of expressed genes that are of less interest or that might complicate analysis (e.g., some or all of the sequences associated with housekeeping genes can be eliminated from REP data).

TEPs can be generated in a manner similar to REPs, e.g., by hybridizing a test sample to an array having a selected set of polynucleotides, particularly a selected set of differentially expressed polynucleotides, acquiring the hybridization data from the array, and storing the data in a format that allows for ready comparison of the TEP with a REP. The REP and TEP to be used in a comparison can be generated simultaneously, or the TEP can be compared to previously generated and stored REPs.

In one embodiment of the invention, comparison of a TEP with a REP involves hybridizing a test sample with a reference array, where the reference array has one or more reference sequences for use in hybridization with a sample. The reference sequences include all, at least one of, or any subset of the differentially expressed polynucleotides described herein. Hybridization data for the test sample is acquired, the data normalized, and the produced TEP compared with a REP generated using an array having the same or similar selected set of differentially expressed polynucleotides. Probes that correspond to sequences differentially expressed between the two samples will show decreased or increased hybridization efficiency for one of the samples relative to the other.

Methods for collection of data from hybridization of samples with a reference arrays are well known in the art. For example, the polynucleotides of the reference and test samples can be generated using a detectable fluorescent label, and hybridization of the polynucleotides in the samples detected by scanning the microarrays for the presence of the detectable label using, for example, a microscope and light source for directing light at a substrate. A photon counter detects fluorescence from the substrate, while an x-y translation stage varies the location of the substrate. A confocal detection device that can be used in the subject methods is described in U.S. Patent No. 5,631,734. A scanning laser microscope is described in Shalon et al., Genome Res. (1996) 6:639. A scan, using the appropriate excitation line, is performed for each fluorophore used. The digital images generated from the scan are then combined for subsequent analysis. For any particular array element, the ratio of the fluorescent signal from one sample (e.g., a test sample) is compared to the fluorescent signal from another sample (e.g., a reference sample), and the relative signal intensity determined.

Methods for analyzing the data collected from hybridization to arrays are well known in the art. For example, where detection of hybridization involves a fluorescent label, data analysis can include the steps of determining fluorescent intensity as a function of substrate position from the data collected, removing outliers, *i.e.*, data deviating from a predetermined statistical distribution, and calculating the relative binding affinity of the targets from the remaining data. The resulting data can be displayed as an image with the intensity in each region varying according to the binding affinity between targets and probes.

In general, the test sample is classified as having a gene expression profile corresponding to that associated with a disease or non-disease state by comparing the TEP generated from the test sample to one or more REPs generated from reference samples (e.g., from samples associated with cancer or specific stages of cancer, dysplasia, samples affected by a disease other than cancer, normal samples, etc.). The criteria for a match or a substantial match between a TEP and a REP include expression of the same or substantially the same set of reference genes, as well as expression of these reference genes at substantially the same levels (e.g., no significant difference between the samples for a signal associated with a selected reference sequence after normalization of the samples, or at least no greater than about 25% to about 40% difference in signal strength for a given reference sequence. In general, a pattern match between a TEP and a REP includes a match in expression, preferably a match in qualitative or quantitative expression level, of at least one of, all or any subset of the differentially expressed genes of the invention.

Pattern matching can be performed manually, or can be performed using a computer program. Methods for preparation of substrate matrices (e.g., arrays), design of oligonucleotides for use with such matrices, labeling of probes, hybridization conditions, scanning of hybridized matrices, and analysis of patterns generated, including comparison analysis, are described in, for example, U.S. Patent No. 5,800,992.

Diagnosis, Prognosis and Management of Cancer

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The polynucleotides of the invention and their gene products are of particular interest as genetic or biochemical markers (e.g., in blood or tissues) that will detect the earliest changes along the carcinogenesis pathway and/or to monitor the efficacy of various therapies and preventive interventions. For example, the level of expression of certain polynucleotides can be indicative of a poorer prognosis, and

therefore warrant more aggressive chemo- or radio-therapy for a patient or vice versa. The correlation of novel surrogate tumor specific features with response to treatment and outcome in patients can define prognostic indicators that allow the design of tailored therapy based on the molecular profile of the tumor. These therapies include antibody targeting and gene therapy. Determining expression of certain polynucleotides and comparison of a patients profile with known expression in normal tissue and variants of the disease allows a determination of the best possible treatment for a patient, both in terms of specificity of treatment and in terms of comfort level of the patient. Surrogate tumor markers, such as polynucleotide expression, can also be used to better classify, and thus diagnose and treat, different forms and disease states of cancer. Two classifications widely used in oncology that can benefit from identification of the expression levels of the polynucleotides of the invention are staging of the cancerous disorder, and grading the nature of the cancerous tissue.

The polynucleotides of the invention can be useful to monitor patients having or susceptible to cancer to detect potentially malignant events at a molecular level before they are detectable at a gross morphological level. Furthermore, a polynucleotide of the invention identified as important for one type of cancer can also have implications for development or risk of development of other types of cancer, e.g., where a polynucleotide is differentially expressed across various cancer types. Thus, for example, expression of a polynucleotide that has clinical implications for metastatic colon cancer can also have clinical implications for stomach cancer or endometrial cancer.

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Staging. Staging is a process used by physicians to describe how advanced the cancerous state is in a patient. Generally, if a cancer is only detectable in the area of the primary lesion without having spread to any lymph nodes it is called Stage I. If it has spread only to the closest lymph nodes, it is called Stage II. In Stage III, the cancer has generally spread to the lymph nodes in near proximity to the site of the primary lesion. Cancers that have spread to a distant part of the body, such as the liver, bone, brain or other site, are Stage IV, the most advanced stage.

The polynucleotides of the invention can facilitate fine-tuning of the staging process by identifying markers for the aggresivity of a cancer, e.g., the metastatic potential, as well as the presence in different areas of the body. Thus, a Stage II cancer with a polynucleotide signifying a high metastatic potential cancer can be used to change a borderline Stage II tumor to a Stage III tumor, justifying more aggressive

therapy. Conversely, the presence of a polynucleotide signifying a lower metastatic potential allows more conservative staging of a tumor.

Grading of cancers. Grade is a term used to describe how closely a tumor resembles normal tissue of its same type. The microscopic appearance of a tumor is used to identify tumor grade based on parameters such as cell morphology, cellular organization, and other markers of differentiation. As a general rule, the grade of a tumor corresponds to its rate of growth or aggressiveness, with undifferentiated or high-grade tumors being more aggressive than well differentiated or low-grade tumors. The following guidelines are generally used for grading tumors: 1) GX Grade cannot be assessed; 2) G1 Well differentiated; G2 Moderately well differentiated; 3) G3 Poorly differentiated; 4) G4 Undifferentiated. The polynucleotides of the invention can be especially valuable in determining the grade of the tumor, as they not only can aid in determining the differentiation status of the cells of a tumor, they can also identify factors other than differentiation that are valuable in determining the aggressivity of a tumor, such as metastatic potential.

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Detection of lung cancer. The polynucleotides of the invention can be used to detect lung cancer in a subject. Although there are more than a dozen different kinds of lung cancer, the two main types of lung cancer are small cell and nonsmall cell, which encompass about 90% of all lung cancer cases. Small cell carcinoma (also called oat cell carcinoma) usually starts in one of the larger bronchial tubes, grows fairly rapidly, and is likely to be large by the time of diagnosis. Nonsmall cell lung cancer (NSCLC) is made up of three general subtypes of lung cancer. Epidermoid carcinoma (also called squamous cell carcinoma) usually starts in one of the larger bronchial tubes and grows relatively slowly. The size of these tumors can range from very small to quite large. Adenocarcinoma starts growing near the outside surface of the lung and can vary in both size and growth rate. Some slowly growing adenocarcinomas are described as alveolar cell cancer. Large cell carcinoma starts near the surface of the lung, grows rapidly, and the growth is usually fairly large when diagnosed. Other less common forms of lung cancer are carcinoid, cylindroma, mucoepidermoid, and malignant mesothelioma.

The polynucleotides of the invention, e.g., polynucleotides differentially expressed in normal cells versus cancerous lung cells (e.g., tumor cells of high or low metastatic potential) or between types of cancerous lung cells (e.g., high metastatic versus low metastatic), can be used to distinguish types of lung cancer as well as identifying traits specific to a certain patient's cancer and selecting an appropriate

therapy. For example, if the patient's biopsy expresses a polynucleotide that is associated with a low metastatic potential, it may justify leaving a larger portion of the patient's lung in surgery to remove the lesion. Alternatively, a smaller lesion with expression of a polynucleotide that is associated with high metastatic potential may justify a more radical removal of lung tissue and/or the surrounding lymph nodes, even if no metastasis can be identified through pathological examination.

The majority of breast cancers are Detection of breast cancer. adenocarcinomas subtypes, which can be summarized as follows: 1) ductal carcinoma in situ (DCIS), including comedocarcinoma; 2) infiltrating (or invasive) ductal carcinoma (IDC); 3) lobular carcinoma in situ (LCIS); 4) infiltrating (or invasive) lobular carcinoma (ILC); 5) inflammatory breast cancer; 6) medullary carcinoma; 7) mucinous carcinoma; 8) Paget's disease of the nipple; 9) Phyllodes tumor; and 10) tubular carcinoma.

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The expression of polynucleotides of the invention can be used in the diagnosis and management of breast cancer, as well as to distinguish between types of breast cancer. Detection of breast cancer can be determined using expression levels of any of the appropriate polynucleotides of the invention, either alone or in combination. Determination of the aggressive nature and/or the metastatic potential of a breast cancer can also be determined by comparing levels of one or more polynucleotides of the invention and comparing levels of another sequence known to vary in cancerous tissue, e.g., ER expression. In addition, development of breast cancer can be detected by examining the ratio of expression of a differentially expressed polynucleotide to the levels of steroid hormones (e.g., testosterone or estrogen) or to other hormones (e.g., growth hormone, insulin). Thus expression of specific marker polynucleotides can be used to discriminate between normal and cancerous breast tissue, to discriminate 25 between breast cancers with different cells of origin, to discriminate between breast cancers with different potential metastatic rates, etc.

The polynucleotides of the invention Detection of colon cancer. exhibiting the appropriate expression pattern can be used to detect colon cancer in a subject. Colorectal cancer is one of the most common neoplasms in humans and perhaps the most frequent form of hereditary neoplasia. Prevention and early detection are key factors in controlling and curing colorectal cancer. Colorectal cancer begins as polyps, which are small, benign growths of cells that form on the inner lining of the colon. Over a period of several years, some of these polyps accumulate additional mutations and become cancerous. Multiple familial colorectal cancer disorders have

been identified, which are summarized as follows: 1) Familial adenomatous polyposis (FAP); 2) Gardner's syndrome; 3) Hereditary nonpolyposis colon cancer (HNPCC); and 4) Familial colorectal cancer in Ashkenazi Jews. The expression of appropriate polynucleotides of the invention can be used in the diagnosis, prognosis and management of colorectal cancer. Detection of colon cancer can be determined using expression levels of any of these sequences alone or in combination with the levels of expression. Determination of the aggressive nature and/or the metastatic potential of a colon cancer can be determined by comparing levels of one or more polynucleotides of the invention and comparing total levels of another sequence known to vary in cancerous tissue, e.g., expression of p53, DCC ras, lor FAP (see, e.g., Fearon ER, et al., Cell (1990) 61(5):759; Hamilton SR et al., Cancer (1993) 72:957; Bodmer W, et al., Nat Genet. (1994) 4(3):217; Fearon ER, Ann N Y Acad Sci. (1995) 768:101). For example, development of colon cancer can be detected by examining the ratio of any of the polynucleotides of the invention to the levels of oncogenes (e.g., ras) or tumor Thus expression of specific marker suppressor genes (e.g., FAP or p53). polynucleotides can be used to discriminate between normal and cancerous colon tissue, to discriminate between colon cancers with different cells of origin, to discriminate between colon cancers with different potential metastatic rates, etc.

Use of Polynucleotides to Screen for Peptide Analogs and Antagonists

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Polypeptides encoded by the instant polynucleotides and corresponding full length genes can be used to screen peptide libraries to identify binding partners, such as receptors, from among the encoded polypeptides. Peptide libraries can be synthesized according to methods known in the art (see, e.g., U.S. Patent No. 5,010,175, and WO 91/17823). Agonists or antagonists of the polypeptides if the invention can be screened using any available method known in the art, such as signal transduction, antibody binding, receptor binding, mitogenic assays, chemotaxis assays, etc. The assay conditions ideally should resemble the conditions under which the native activity is exhibited in vivo, that is, under physiologic pH, temperature, and ionic strength. Suitable agonists or antagonists will exhibit strong inhibition or enhancement of the native activity at concentrations that do not cause toxic side effects in the subject. Agonists or antagonists that compete for binding to the native polypeptide can require concentrations equal to or greater than the native concentration, while inhibitors capable of binding irreversibly to the polypeptide can be added in concentrations on the order of the native concentration.

Such screening and experimentation can lead to identification of a novel polypeptide binding partner, such as a receptor, encoded by a gene or a cDNA corresponding to a polynucleotide of the invention, and at least one peptide agonist or antagonist of the novel binding partner. Such agonists and antagonists can be used to modulate, enhance, or inhibit receptor function in cells to which the receptor is native, or in cells that possess the receptor as a result of genetic engineering. Further, if the novel receptor shares biologically important characteristics with a known receptor, information about agonist/antagonist binding can facilitate development of improved agonists/antagonists of the known receptor.

10 Pharmaceutical Compositions and Therapeutic Uses

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can invention the Pharmaceutical compositions of polypeptides, antibodies, or polynucleotides (including antisense nucleotides and ribozymes) of the claimed invention in a therapeutically effective amount. The term "therapeutically effective amount" as used herein refers to an amount of a therapeutic agent to treat, ameliorate, or prevent a desired disease or condition, or to exhibit a detectable therapeutic or preventative effect. The effect can be detected by, for example, chemical markers or antigen levels. Therapeutic effects also include reduction in physical symptoms, such as decreased body temperature. The precise effective amount for a subject will depend upon the subject's size and health, the nature and extent of the condition, and the therapeutics or combination of therapeutics selected for administration. Thus, it is not useful to specify an exact effective amount in advance. However, the effective amount for a given situation is determined by routine experimentation and is within the judgment of the clinician. For purposes of the present invention, an effective dose will generally be from about 0.01 mg/ kg to 50 mg/kg or 0.05 mg/kg to about 10 mg/kg of the DNA constructs in the individual to which it is administered.

A pharmaceutical composition can also contain a pharmaceutically acceptable carrier. The term "pharmaceutically acceptable carrier" refers to a carrier for administration of a therapeutic agent, such as antibodies or a polypeptide, genes, and other therapeutic agents. The term refers to any pharmaceutical carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition, and which can be administered without undue toxicity. Suitable carriers can be large, slowly metabolized macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers,

and inactive virus particles. Such carriers are well known to those of ordinary skill in the art. Pharmaceutically acceptable carriers in therapeutic compositions can include liquids such as water, saline, glycerol and ethanol. Auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, can also be present in such vehicles. Typically, the therapeutic compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection can also be prepared. Liposomes are included within the definition of a pharmaceutically acceptable carrier. Pharmaceutically acceptable salts can also be present in the pharmaceutical composition, e.g., mineral acid salts such as hydrochlorides, hydrobromides, phosphates, sulfates, and the like; and the salts of organic acids such as acetates, propionates, malonates, benzoates, and the like. A thorough discussion of pharmaceutically acceptable excipients is available in Remington's Pharmaceutical Sciences (Mack Pub. Co., New Jersey, 1991).

Delivery Methods. Once formulated, the compositions of the invention can be (1) administered directly to the subject (e.g., as polynucleotide or polypeptides); or (2) delivered ex vivo, to cells derived from the subject (e.g., as in ex vivo gene therapy). Direct delivery of the compositions will generally be accomplished by parenteral injection, e.g., subcutaneously, intraperitoneally, intravenously or intramuscularly, intratumoral or to the interstitial space of a tissue. Other modes of administration include oral and pulmonary administration, suppositories, and transdermal applications, needles, and gene guns or hyposprays. Dosage treatment can be a single dose schedule or a multiple dose schedule.

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Methods for the ex vivo delivery and reimplantation of transformed cells into a subject are known in the art and described in e.g., International Publication No. WO 93/14778. Examples of cells useful in ex vivo applications include, for example, stem cells, particularly hematopoetic, lymph cells, macrophages, dendritic cells, or tumor cells. Generally, delivery of nucleic acids for both ex vivo and in vitro applications can be accomplished by, for example, dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei, all well known in the art.

Once a gene corresponding to a polynucleotide of the invention has been found to correlate with a proliferative disorder, such as neoplasia, dysplasia, and hyperplasia, the disorder can be amenable to treatment by administration of a

therapeutic agent based on the provided polynucleotide, corresponding polypeptide or other corresponding molecule (e.g., antisense, ribozyme, etc.).

The dose and the means of administration of the inventive pharmaceutical compositions are determined based on the specific qualities of the therapeutic composition, the condition, age, and weight of the patient, the progression For example, administration of of the disease, and other relevant factors. polynucleotide therapeutic compositions agents of the invention includes local or systemic administration, including injection, oral administration, particle gun or catheterized administration, and topical administration. Preferably, the therapeutic polynucleotide composition contains an expression construct comprising a promoter operably linked to a polynucleotide of at least 12, 22, 25, 30, or 35 contiguous nt of the polynucleotide disclosed herein. Various methods can be used to administer the therapeutic composition directly to a specific site in the body. For example, a small metastatic lesion is located and the therapeutic composition injected several times in several different locations within the body of tumor. Alternatively, arteries which serve a tumor are identified, and the therapeutic composition injected into such an artery, in order to deliver the composition directly into the tumor. A tumor that has a necrotic center is aspirated and the composition injected directly into the now empty center of the tumor. The antisense composition is directly administered to the surface of the tumor, for example, by topical application of the composition. X-ray imaging is used to assist in certain of the above delivery methods.

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Receptor-mediated targeted delivery of therapeutic compositions containing an antisense polynucleotide, subgenomic polynucleotides, or antibodies to specific tissues can also be used. Receptor-mediated DNA delivery techniques are described in, for example, Findeis et al., *Trends Biotechnol.* (1993) 11:202; Chiou et al., *Gene Therapeutics: Methods And Applications Of Direct Gene Transfer* (J.A. Wolff, ed.) (1994); Wu et al., *J. Biol. Chem.* (1988) 263:621; Wu et al., *J. Biol. Chem.* (1994) 269:542; Zenke et al., *Proc. Natl. Acad. Sci.* (USA) (1990) 87:3655; Wu et al., *J. Biol. Chem.* (1991) 266:338. Therapeutic compositions containing a polynucleotide are administered in a range of about 100 ng to about 200 mg of DNA for local administration in a gene therapy protocol. Concentration ranges of about 500 ng to about 50 mg, about 1 mg to about 2 mg, about 5 mg to about 500 mg, and about 20 mg to about 100 mg of DNA can also be used during a gene therapy protocol. Factors such as method of action (e.g., for enhancing or inhibiting levels of the encoded gene product) and efficacy of transformation and expression are considerations which will

affect the dosage required for ultimate efficacy of the antisense subgenomic polynucleotides. Where greater expression is desired over a larger area of tissue, larger amounts of antisense subgenomic polynucleotides or the same amounts readministered in a successive protocol of administrations, or several administrations to different adjacent or close tissue portions of, for example, a tumor site, may be required to effect a positive therapeutic outcome. In all cases, routine experimentation in clinical trials will determine specific ranges for optimal therapeutic effect. For polynucleotide-related genes encoding polypeptides or proteins with anti-inflammatory activity, suitable use, doses, and administration are described in U.S. Patent No. 5,654,173.

The therapeutic polynucleotides and polypeptides of the present invention can be delivered using gene delivery vehicles. The gene delivery vehicle can be of viral or non-viral origin (see generally, Jolly, Cancer Gene Therapy (1994) 1:51; Kimura, Human Gene Therapy (1994) 5:845; Connelly, Human Gene Therapy (1995) 1:185; and Kaplitt, Nature Genetics (1994) 6:148). Expression of such coding sequences can be induced using endogenous mammalian or heterologous promoters. Expression of the coding sequence can be either constitutive or regulated.

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Viral-based vectors for delivery of a desired polynucleotide and expression in a desired cell are well known in the art. Exemplary viral-based vehicles include, but are not limited to, recombinant retroviruses (see, e.g., WO 90/07936; WO 94/03622; WO 93/25698; WO 93/25234; U.S. Patent No. 5, 219,740; WO 93/11230; WO 93/10218; U.S. Patent No. 4,777,127; GB Patent No. 2,200,651; EP 0 345 242; and WO 91/02805), alphavirus-based vectors (e.g., Sindbis virus vectors, Semliki forest virus (ATCC VR-67; ATCC VR-1247), Ross River virus (ATCC VR-373; ATCC VR-1246) and Venezuelan equine encephalitis virus (ATCC VR-923; ATCC VR-1250; ATCC VR 1249; ATCC VR-532), and adeno-associated virus (AAV) vectors (see, e.g., WO 94/12649, WO 93/03769; WO 93/19191; WO 94/28938; WO 95/11984 and WO 95/00655). Administration of DNA linked to killed adenovirus as described in Curiel, Hum. Gene Ther. (1992) 3:147 can also be employed.

Non-viral delivery vehicles and methods can also be employed, including, but not limited to, polycationic condensed DNA linked or unlinked to killed adenovirus alone (see, e.g., Curiel, Hum. Gene Ther. (1992) 3:147); ligand-linked DNA(see, e.g., Wu, J. Biol. Chem. 264:16985 (1989)); eukaryotic cell delivery vehicles cells (see, e.g., U.S. Patent No. 5,814,482; WO 95/07994; WO 96/17072; WO 95/30763; and WO 97/42338) and nucleic charge neutralization or fusion with cell membranes. Naked DNA can also be employed. Exemplary naked DNA introduction

methods are described in WO 90/11092 and U.S. Patent No. 5,580,859. Liposomes that can act as gene delivery vehicles are described in U.S. Patent No. 5,422,120; WO 95/13796; WO 94/23697; WO 91/14445; and EP 0524968. Additional approaches are described in Philip, Mol. Cell Biol. 14:2411 (1994), and in Woffendin, Proc. Natl. Acad. Sci. (1994) 91:1581.

Further non-viral delivery suitable for use includes mechanical delivery systems such as the approach described in Woffendin et al., *Proc. Natl. Acad. Sci. USA* 91(24):11581 (1994). Moreover, the coding sequence and the product of expression of such can be delivered through deposition of photopolymerized hydrogel materials or use of ionizing radiation (see, *e.g.*, U.S. Patent No. 5,206,152 and WO 92/11033). Other conventional methods for gene delivery that can be used for delivery of the coding sequence include, for example, use of hand-held gene transfer particle gun (see, *e.g.*, U.S. Patent No. 5,149,655); use of ionizing radiation for activating transferred gene (see, *e.g.*, U.S. Patent No. 5,206,152 and WO 92/11033).

The present invention will now be illustrated by reference to the following examples which set forth particularly advantageous embodiments. However, it should be noted that these embodiments are illustrative and are not to be construed as restricting the invention in any way.

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WO 01/02568

EXAMPLES

EXAMPLE 1

SOURCE OF BIOLOGICAL MATERIALS AND OVERVIEW OF NOVEL POLYNUCLEOTIDES

EXPRESSED BY THE BIOLOGICAL MATERIALS

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Cell lines and human normal and tumor tissue were used to construct cDNA libraries from mRNA isolated from the cells and tissues. Most sequences were about 275-300 nucleotides in length. The cells lines include Km12L4-A cell line, a high metastatic colon cancer cell line (Morika, W. A. K. et al., Cancer Research (1988) 48:6863). The KM12L4-A cell line is derived from the KM12C cell line. The KM12C cell line, which is poorly metastatic (low metastatic) was established in culture from a Dukes' stage B2 surgical specimen (Morikawa et al. Cancer Res. (1988) 48:6863). The KML4-A is a highly metastatic subline derived from KM12C (Yeatman et al. Nucl. Acids. Res. (1995) 23:4007; Bao-Ling et al. Proc. Annu. Meet. Am. Assoc. Cancer. Res. (1995) 21:3269). The KM12C and KM12C-derived cell lines (e.g., KM12L4, KM12L4-A, etc.) are well-recognized in the art as model cell lines for the study of colon cancer (see, e.g., Moriakawa et al., supra; Radinsky et al. Clin. Cancer Res. (1995) 1:19; Yeatman et al., (1995) supra; Yeatman et al., Clin. Exp. Metastasis (1996) 14:246). These and other cell lines and tissue are described in Table 6.

The sequences of the isolated polynucleotides were first masked to eliminate low complexity sequences using the XBLAST masking program (Claverie "Effective Large-Scale Sequence Similarity Searches," In: Computer Methods for Macromolecular Sequence Analysis, Doolittle, ed., Meth. Enzymol. 266:212-227 Academic Press, NY, NY (1996); see particularly Claverie, in "Automated DNA Sequencing and Analysis Techniques" Adams et al., eds., Chap. 36, p. 267 Academic Press, San Diego, 1994 and Claverie et al. Comput. Chem. (1993) 17:191). Generally, masking does not influence the final search results, except to eliminate sequences of relative little interest due to their low complexity, and to eliminate multiple "hits" based on similarity to repetitive regions common to multiple sequences, e.g., Alu repeats. The sequences remaining after masking were then used in a BLASTN vs. Genbank search; sequences that exhibited greater than 70% overlap, 99% identity, and a p value of less than 1 x 10⁴⁰ were discarded. Sequences from this search also were discarded if the inclusive parameters were met, but the sequence was ribosomal or vector-derived.

The resulting sequences from the previous search were classified into three groups (1, 2 and 3 below) and searched in a BLASTX vs. NRP (non-redundant proteins) database search: (1) unknown (no hits in the Genbank search), (2) weak similarity (greater than 45% identity and p value of less than 1 x 10^{-5}), and (3) high similarity (greater than 60% overlap, greater than 80% identity, and p value less than 1 x 10^{-5}). Sequences having greater than 70% overlap, greater than 99% identity, and p value of less than 1 x 10^{-40} were discarded.

The remaining sequences were classified as unknown (no hits), weak similarity, and high similarity (parameters as above). Two searches were performed on these sequences. First, a BLAST vs. EST database search was performed and sequences with greater than 99% overlap, greater than 99% similarity and a p value of less than 1 x 10⁻⁴⁰ were discarded. Sequences with a p value of less than 1 x 10⁻⁶⁵ when compared to a database sequence of human origin were also excluded. Second, a BLASTN vs. Patent GeneSeq database was performed and sequences having greater than 99% identity, p value less than 1 x 10⁻⁴⁰, and greater than 99% overlap were discarded.

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The remaining sequences were subjected to screening using other rules and redundancies in the dataset. Sequences with a p value of less than 1 x 10⁻¹¹¹ in relation to a database sequence of human origin were specifically excluded. The final result provided the 3351 sequences listed in the accompanying Sequence Listing. Each identified polynucleotide represents sequence from at least a partial mRNA transcript. Polynucleotides that were determined to be novel were assigned a sequence identification number.

The novel polynucleotides were assigned sequence identification numbers SEQ 1D NOs:1-3351. The first 1847 DNA sequences corresponding to the novel polynucleotides are provided in the Sequence Listing in Table 1. DNA sequences corresponding to the novel polynucleotides of SEQ 1D NOs:1848-3351 are provided in the Sequence Listing in Table 2. The DNA sequences of Table 2, while numbered SEQ 1D 1-1504, correspond to SEQ 1D NOs:1848-3351 in the Sequence Listing, e.g., Table 2 SEQ 1D 1 is SEQ 1D NO:1848, Table 2 SEQ 1D 2 is SEQ 1D NO:1849, etc. Each DNA sequence in Table 4 is uniquely identified by a number that is 1847 less than its SEQ 1D NO in the Sequence Listing. Tables 1 and 2 provide: 1) the SEQ 1D NO assigned to cach sequence for use in the present specification or a corresponding number; 2) the sequence name used as an internal identifier of the sequence; 3) the name assigned to the clone from which the

sequence was isolated; and 4) the number of the cluster to which the sequence is assigned (Cluster ID; where the cluster ID is 0, the sequence was not assigned to any cluster).

Because the provided polynucleotides represent partial mRNA transcripts, two or more polynucleotides of the invention may represent different regions of the same mRNA transcript and the same gene. Thus, if two or more SEQ ID NOs: are identified as belonging to the same clone, then either sequence can be used to obtain the full-length mRNA or gene.

EXAMPLE 2

RESULTS OF PUBLIC DATABASE SEARCH TO IDENTIFY FUNCTION OF GENE PRODUCTS

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SEQ ID NOs:1-3351 were translated in all three reading frames to determine the best alignment with the individual sequences. These amino acid sequences and nucleotide sequences are referred to, generally, as query sequences, which are aligned with the individual sequences. Query and individual sequences were aligned using the BLAST programs, available over the world wide web at http://www.ncbi.nlm.nih.gov/BLAST/. Again the sequences were masked to various extents to prevent searching of repetitive sequences or poly-A sequences, using the XBLAST program for masking low complexity as described above in Example 1.

Tables 3 and 4 (inserted before the claims) show the results of the alignments. Table 3 contains alignment information for SEQ ID NOs:1-1847 and Table 4 contains alignment information for SEQ ID NOs:1848-3351. The DNA sequences of Table 4, while numbered SEQ ID 1-1504, correspond to SEQ ID NOs:1848-3351. Each DNA sequence in Table 4 is uniquely identified by a number that is 1847 less than its SEQ ID NO. Tables 3 and 4 refer to each sequence by its SEQ ID NO or a corresponding number, the accession numbers and descriptions of nearest neighbors from the Genbank and Non-Redundant Protein searches, and the p values of the search results.

For each of SEQ ID NOs:1-1847, the best alignment to a protein or DNA sequence is included in Table 3, and the best alignment for each of SEQ ID NOs:1848-3351 is included in Table 4. The activity of the polypeptide encoded by SEQ ID NOs:1-3351 is the same or similar to the nearest neighbor reported in Table 3 or 4. The accession number of the nearest neighbor is reported, providing a reference to the activities exhibited by the nearest neighbor. The search program and database used for the alignment also are indicated as well as a calculation of the p value.

Full length sequences or fragments of the polynucleotide sequences of the nearest neighbors can be used as probes and primers to identify and isolate the full length sequence of SEQ ID NOs:1-3351. The nearest neighbors can indicate a tissue or cell type to be used to construct a library for the full-length sequences of SEQ ID NOs:1-3351.

EXAMPLE 3 Members of Protein Families

The sequences (SEQ ID NOs:1-3351) were used to conduct a profile scarch as described in the specification above. Several of the polynucleotides of the invention were found to encode polypeptides having characteristics of a polypeptide belonging to a known protein families (and thus represent new members of these protein families) and/or comprising a known functional domain (Table 5). "Start" and "stop" in Table 3 indicate the position within the individual sequences that align with the query sequence having the indicated SEQ ID NO. The direction indicates the orientation of the query sequence with respect to the individual sequence, where forward (for) indicates that the alignment is in the same direction (left to right) as the sequence provided in the Sequence Listing and reverse (rev) indicates that the alignment is with a sequence complementary to the sequence provided in the Sequence

Some polynucleotides exhibited multiple profile hits because, for example, the particular sequence contains overlapping profile regions, and/or the sequence contains two different functional domains. These profile hits are described in more detail below.

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Ank Repeats (ANK). SEQ ID NOs:187, 1268, 1804, 1819, 1830, 1839, 2652, 3015 and 3267 represent polynucleotides encoding an Ank repeat-containing protein. The ankyrin motif is a 33 amino acid sequence named for the protein ankyrin which has 24 tandem 33-amino-acid motifs. Ank repeats were originally identified in the cell-cycle-control protein cdc10 (Breeden et al., Nature (1987) 329:651). Proteins containing ankyrin repeats include ankyrin, myotropin, I-kappaB proteins, cell cycle protein cdc10, the Notch receptor (Matsuno et al., Development (1997) 124(21):4265); G9a (or BAT8) of the class III region of the major histocompatibility complex (Biochem J. 290:811-818, 1993), FABP, GABP, 53BP2, Lin12, glp-1, SW14, and SW16. The functions of the ankyrin repeats are compatible with a role in protein-

protein interactions (Bork, Proteins (1993) 17(4):363; Lambert and Bennet, Eur. J. Biochem. (1993) 211:1; Kerr et al., Current Op. Cell Biol. (1992) 4:496; Bennet et al., J. Biol. Chem. (1980) 255:6424).

ATPases Associated with Various Cellular Activities (ATPases). Sequences within SEQ ID NOs:431, 639, 2135, 2684, 2859, 3197 and 3266 correspond to a sequence that encodes a novel member of the "ATPases Associated with diverse cellular Activities" (AAA) protein family. The AAA protein family is composed of a large number of ATPases that share a conserved region of about 220 amino acids that contains an ATP-binding site (Froehlich et al., J. Cell Biol. (1991) 114:443; Erdmann et al., Cell (1991) 64:499; Peters et al., EMBO J. (1990) 9:1757; Kunau et al., Biochimie (1995)**BioEssays** al., Confalonieri et 75:209-224; (1993)http://yeamob.pci.chemie.uni-tuebingen.de/AAA/Description.html). The proteins that belong to this family either contain one or two AAA domains. In general, the AAA domains in these proteins act as ATP-dependent protein clamps (Confalonieri et al. (1995) BioEssays 17:639). In addition to the ATP-binding 'A' and 'B' motifs, which are located in the N-terminal half of this domain, there is a highly conserved region located in the central part of the domain which was used in the development of the signature pattern. The consensus pattern is: [LIVMT]-x-[LIVMT]-[LIVMF]-x-[GATMC]-[ST]-[NS]-x(4)-[LIVM]-D-x-A-[LIFA]-x-R.

Bromodomain (bromodomain). SEQ ID NO:1814 represents a polynucleotide encoding a polypeptide having a bromodomain region (Haynes et al., 1992, Nucleic Acids Res. 20:2693-2603, Tamkun et al., 1992, Cell 68:561-572, and Tamkun, 1995, Curr. Opin. Genet. Dev. 5:473-477), which is a conserved region of about 70 amino acids. The bromodomain is thought to be involved in protein-protein interactions and may be important for the assembly or activity of multicomponent complexes involved in transcriptional activation. The consensus pattern, which spans a major part of the bromodomain, is: [STANVF]-x(2)-F-x(4)-[DNS]-x(5,7)-[DENQTF]-Y-[HFY]-x(2)- [LIVMFY]-x(3)-[LIVM]-x(4)-[LIVM]-x(6,8)-Y-x(12,13)-[LIVM]-x(2)-N-[SACF]-x(2)-[FY].

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Basic Region Plus Leucine Zipper Transcription Factors (BZIP). SEQ ID NOs:410, 552, 768, 822, 836, 1288, 1365, 1454, 1540, 1549, 1556, 1557, 1563, 1622, 1630, 1704, 1808, 2363, 2424, 3147, 3152, 3158 and 3208 represent polynucleotides encoding a novel member of the family of basic region plus leucine zipper transcription factors. The bZIP superfamily (Hurst, *Protein Prof.* (1995) 2:105; and Ellenberger, *Curr. Opin. Struct. Biol.* (1994) 4:12) of eukaryotic DNA-binding

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transcription factors encompasses proteins that contain a basic region mediating sequence-specific DNA-binding followed by a leucine zipper required for dimerization. The consensus pattern for this protein family is: [KR]-x(1,3)-[RKSAQ]-N-x(2)-[SAQ](2)-x-[RKTAENQ]-x-R-x-[RK].

EF Hand (EFhand). SEQ ID NOs:820, 1755 and 3285 correspond to polynucleotides encoding a novel protein in the family of EF-hand proteins. Many calcium-binding proteins belong to the same evolutionary family and share a type of calcium-binding domain known as the EF-hand (Kawasaki et al., *Protein. Prof.* (1995) 2:305-490). This type of domain consists of a twelve residue loop flanked on both sides by a twelve residue alpha-helical domain. In an EF-hand loop the calcium ion is coordinated in a pentagonal bipyramidal configuration. The six residues involved in the binding are in positions 1, 3, 5, 7, 9 and 12; these residues are denoted by X, Y, Z, -Y, -X and -Z. The invariant Glu or Asp at position 12 provides two oxygens for liganding Ca (bidentate ligand). The consensus pattern includes the complete EF-hand loop as well as the first residue which follows the loop and which seem to always be hydrophobic:

D-x-[DNS]-{ILVFYW}-[DENSTG]-[DNQGHRK]-{GP}-[LIVMC]-[DENQSTAGC]-x(2)-[DE]-[LIVMFYW].

Ets Domain (Ets Nterm). SEQ ID NO:1811 represents a polynucleotide encoding a polypeptide with N-terminal homology in ETS domain. Proteins of this family contain a conserved domain, the "ETS-domain," that is involved in DNA binding. The domain appears to recognize purine-rich sequences; it is about 85 to 90 amino acids in length, and is rich in aromatic and positively charged residues (Wasylyk, et al., Eur. J. Biochem. (1993) 211:718). The ets gene family encodes a novel class of DNA-binding proteins, each of which binds a specific DNA sequence and comprises an ets domain that specifically interacts with sequences containing the common core trinucleotide sequence GGA. In addition to an ets domain, native ets proteins comprise other sequences which can modulate the biological specificity of the protein. Ets genes and proteins are involved in a variety of essential biological processes including cell growth, differentiation and development, and three members are implicated in oncogenic process.

G-Protein Alpha Subunit (G-alpha). SEQ ID NO:1846 represents a polynucleotide encoding a novel polypeptide of the G-protein alpha subunit family. Guanine nucleotide binding proteins (G-proteins) are a family of membrane-associated proteins that couple extracellularly-activated integral-membrane receptors to intracellular effectors, such as ion channels and enzymes that vary the concentration of

second messenger molecules. G-proteins are composed of 3 subunits (alpha, beta and gamma) which, in the resting state, associate as a trimer at the inner face of the plasma membrane. The alpha subunit binds GTP and exhibits GTPase activity. G-protein alpha subunits are 350-400 amino acids in length and have molecular weights in the range 40-45 kDa. Seventeen distinct types of alpha subunit have been identified in mammals, and fall into 4 main groups on the basis of both sequence similarity and function: alphas, alpha-q, alpha-i and alpha-12 (Simon et al., Science (1993) 252:802). They are often N-terminally acylated, usually with myristate and/or palmitoylate, and these fatty acid modifications can be important for membrane association and high- affinity interactions with other proteins.

10 Helicases conserved C-terminal domain (helicase C). NOs:1496, 2826 and 2871 represent polynucleotides encoding novel members of the DEAD/H helicase family. A number of eukaryotic and prokaryotic proteins have been characterized (Schmid S.R., et al., Mol. Microbiol. (1992) 6:283; Linder P., et al., Nature (1989) 337:121; Wassarman D.A., et al., Nature (1991) 349:463) on the basis of their structural similarity. All are involved in ATP-dependent, nucleic-acid unwinding. 15 All DEAD box family members of the above proteins share a number of conserved sequence motifs, some of which are specific to the DEAD family while others are shared by other ATP-binding proteins or by proteins belonging to the helicases 'superfamily' (Hodgman T.C., Nature (1988) 333:22 and Nature (1988) 333:578 (Errata). One of these motifs, called the "D-E-A-D-box", represents a special version of 20 the B motif of ATP-binding proteins. Some other proteins belong to a subfamily which have His instead of the second Asp and are thus said to be "D-E-A-H-box" proteins (Wassarman D.A., et al., Nature (1991) 349:463; Harosh I., et al., Nucleic Acids Res. (1991) 19:6331; Koonin E.V. et al., J. Gen. Virol. (1992) 73:989. The following signature patterns are used to identify members of both subfamilies: 1) [LIVMF](2)-D-E-A-D-[RKEN]-x-[LIVMFYGSTN]; and 2) [GSAH]-x-[LIVMF](3)-D-E-[ALIV]-H-[NECR].

Homeobox domain (homeobox). SEQ ID NOs:1676, 1820 and 1821 represent polynucleotides encoding proteins having a homeobox domain. The homeobox is a protein domain of 60 amino acids (Gehring In: Guidebook to the Homeobox Genes, Duboule D., Ed., pp. 1-10, Oxford University Press, Oxford, (1994); Buerglin In: Guidebook to the Homeobox Genes, pp25-72, Oxford University Press, Oxford, (1994); Gehring, Trends Biochem. Sci. (1992) 17:277-280; Gehring et al., Annu. Rev. Genet. (1986) 20:147-173; Schofield, Trends Neurosci. (1987) 10:3-6) first

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identified in a number of Drosophila homeotic and segmentation proteins. It is extremely well conserved in many other animals, including vertebrates. This domain binds DNA through a helix-turn-helix type of structure. Several proteins that contain a homeobox domain play an important role in development. Most of these proteins are sequence-specific DNA-binding transcription factors. The homeobox domain is also very similar to a region of the yeast mating type proteins. These are sequence-specific DNA-binding proteins that act as master switches in yeast differentiation by controlling gene expression in a cell type-specific fashion.

A schematic representation of the homeobox domain is shown below.

The helix-turn-helix region is shown by the symbols 'H' (for helix), and 't' (for turn).

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The pattern detects homeobox sequences 24 residues long and spans positions 34 to 57 of the homeobox domain. The consensus pattern is as follows: [LIVMFYG]-[ASLVR]-x(2)-[LIVMSTACN]-x-[LIVM]-x(4)-[LIV]-[RKNQESTAIY]-[LIVFSTNKH]-W-[FYVC]-x-[NDQTAH]-x(5)-[RKNAIMW].

MAP kinase kinase (mkk). SEQ ID NOs:29, 31, 196, 3175, 3190 and 3281 represent novel members of the MAP kinase kinase family. MAP kinases (MAPK) are involved in signal transduction, and are important in cell cycle and cell 20 growth controls. The MAP kinase kinases (MAPKK) are dual-specificity protein kinases which phosphorylate and activate MAP kinases. MAPKK homologues have Moreover, the been found in yeast, invertebrates, amphibians, and mammals. MAPKK/MAPK phosphorylation switch constitutes a basic module activated in distinct pathways in yeast and in vertebrates. MAPKKs are essential transducers through which signals must pass before reaching the nucleus. For review, see, e.g., Biologique Biol Cell (1993) 79:193-207; Nishida et al., Trends Biochem Sci (1993) 18:128-31; Ruderman, Curr Opin Cell Biol (1993) 5:207-13; Dhanasekaran et al., Oncogene (1998) 17:1447-55; Kiefer et al., Biochem Soc Trans (1997) 25:491-8; and Hill, Cell Signal 30 (1996) 8:533-44.

Protein Kinase (protkinase). SEQ ID NOs:1157, 1478, 1496, 2286, 2969 and 3190 represent polynucleotides encoding protein kinases. Protein kinases catalyze phosphorylation of proteins in a variety of pathways, and are implicated in cancer. Eukaryotic protein kinases (Hanks S.K., et al., FASEB J. (1995) 9:576; Hunter T., Meth. Enzymol. (1991) 200:38; Hanks S.K., et al., Meth. Enzymol. (1991) 200:38; Hanks S.K.,

Curr. Opin. Struct. Biol. (1991) 1:369; Hanks S.K. et al., Science (1988) 241:42) are enzymes that belong to a very extensive family of proteins which share a conserved catalytic core common to both serine/threonine and tyrosine protein kinases. There are a number of conserved regions in the catalytic domain of protein kinases. The first region, which is located in the N-terminal extremity of the catalytic domain, is a glycine-rich stretch of residues in the vicinity of a lysine residue, which has been shown to be involved in ATP binding. The second region, which is located in the central part of the catalytic domain, contains a conserved aspartic acid residue which is important for the catalytic activity of the enzyme (Knighton D.R. et al., Science (1991) 253:407).
The protein kinase profile includes two signature patterns for this second region: one specific for serine/threonine kinases and the other for tyrosine kinases. A third profile is based on the alignment in (Hanks S.K. et al., FASEB J. (1995) 9:576) and covers the entire catalytic domain.

The consensus patterns are as follows: 1) [LIV]-G-{P}-G-{P}-15 [FYWMGSTNH]-[SGA]-{PW}-[LIVCAT]-{PD}-x-[GSTACLIVMFY]-x(5,18)[LIVMFYWCSTAR]-[AIVP]-[LIVMFAGCKR]-K, where K binds ATP; 2)
[LIVMFYC]-x-[HY]-x-D-[LIVMFY]-K-x(2)-N-[LIVMFYCT](3), where D is an active site residue; and 3) [LIVMFYC]-x-[HY]-x-D-[LIVMFY]-[RSTAC]-x(2)-N[LIVMFYC], where D is an active site residue.

If a protein analyzed includes two of the above protein kinase signatures, the probability of it being a protein kinase is close to 100%.

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Ras family proteins (ras). SEQ ID NOs:1688 and 3258 represent polynucleotides encoding novel members of the ras family of small GTP/GDP-binding proteins (Valencia et al., 1991, Biochemistry 30:4637-4648). Ras family members generally require a specific guanine nucleotide exchange factor (GEF) and a specific GTPase activating protein (GAP) as stimulators of overall GTPase activity. Among ras-related proteins, the highest degree of sequence conservation is found in four regions that are directly involved in guanine nucleotide binding. The first two constitute most of the phosphate and Mg2+ binding site (PM site) and are located in the first half of the G-domain. The other two regions are involved in guanosine binding and are located in the C-terminal half of the molecule. Motifs and conserved structural features of the ras-related proteins are described in Valencia et al., 1991, Biochemistry 30:4637-4648. A major consensus pattern of ras proteins is: D-T-A-G-Q-E-K-[LF]-G-G-L-R-[DE]-G-Y-Y.

Thioredoxin family active site (Thioredox). SEQ ID NO:1677 represents a polynucleotide encoding a protein having a thioredoxin family active site. Thioredoxins (Holmgren A., Annu. Rev. Biochem. (1985) 54:237; Gleason F.K. et al., FEMS Microbiol. Rev. (1988) 54:271; Holmgren, A. J. Biol. Chem. (1989) 264:13963; Eklund H. et al., Proteins (1991) 11:13) are small proteins of approximately one hundred amino- acid residues which participate in various redox reactions via the reversible oxidation of an active center disulfide bond. They exist in either a reduced form or an oxidized form where the two cysteine residues are linked in an intramolecular disulfide bond. Thioredoxin is present in prokaryotes and eukaryotes and the sequence around the redox-active disulfide bond is well conserved. All PDI contains two or three (ERp72) copies of the thioredoxin domain. The consensus pattern is:

[LIVMF]-[LIVMSTA]-x-[LIVMFYC]-[FYWSTHE]-x(2)-[FYWGTN]-C-[GATPLVE]-[PHYWSTA]-C-x(6)-[LIVMFYWT] (where the two C's form the redoxactive bond).

Trypsin (trypsin). SEQ ID NO:1410 corresponds to a novel serine protease of the trypsin family. The catalytic activity of the serine proteases from the trypsin family is provided by a charge relay system involving an aspartic acid residue hydrogen-bonded to a histidine, which itself is hydrogen-bonded to a serine. The sequences in the vicinity of the active site serine and histidine residues are well conserved in this family of proteases (Brenner S., *Nature* (1988) 334:528). The consensus patterns for this trypsin protein family are: 1) [LIVM]-[ST]-A-[STAG]-H-C, where H is the active site residue; and 2) [DNSTAGC]-[GSTAPIMVQH]-x(2)-G-[DE]-S-G-[GS]-[SAPHV]- [LIVMFYWH]-[LIVMFYSTANQH], where S is the active site residue. All sequences known to belong to this family are detected by the above consensus sequences, except for 18 different proteases which have lost the first conserved glycine. If a protein includes both the serine and the histidine active site signatures, the probability of it being a trypsin family serine protease is 100%.

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WD Domain, G-Beta Repeats (WD domain). SEQ ID NOs:1336, 1380, 1711, 1762, 1909, 2218, 3047, 3108 and 3292 represent novel members of the WD domain/G-beta repeat family. Beta-transducin (G-beta) is one of the three subunits (alpha, beta, and gamma) of the guanine nucleotide-binding proteins (G proteins) which act as intermediaries in the transduction of signals generated by transmembrane receptors (Gilman, Annu. Rev. Biochem. (1987) 56:615). The alpha subunit binds to and hydrolyzes GTP; the functions of the beta and gamma subunits are less clear but they seem to be required for the replacement of GDP by GTP as well as for membrane

anchoring and receptor recognition. In higher eukaryotes, G-beta exists as a small multigene family of highly conserved proteins of about 340 amino acid residues. Structurally, G-beta consists of eight tandem repeats of about 40 residues, each containing a central Trp-Asp motif (this type of repeat is sometimes called a WD-40 repeat). The consensus pattern for the WD domain/G-Beta repeat family is: [LIVMSTAC]-[LIVMFYWSTAGC]-[LIMSTAG]-[LIVMSTAGC]-x(2)-[DN]-x(2)-[LIVMWSTAC]-x-[LIVMFSTAG]-W-[DEN]-[LIVMFSTAGCN].

wnt Family of Developmental Signaling Proteins (Wnt_dev_sign). SEQ ID NO:1538 corresponds to a novel member of the wnt family of developmental signaling proteins. Wnt-1 (previously known as int-1), the seminal member of this family, (Nusse R., Trends Genet. (1988) 4:291) is thought to play a role in intercellular communication and seems to be a signalling molecule important in the development of the central nervous system (CNS). All wnt family proteins share the following features characteristics of secretory proteins: a signal peptide, several potential N-glycosylation sites and 22 conserved cysteines that are probably involved in disulfide bonds. The Wnt proteins seem to adhere to the plasma membrane of the secreting cells and are therefore likely to signal over only few cell diameters. The consensus pattern, which is based upon a highly conserved region including three cysteines, is as follows: C-K-C-H-G-[LIVMT]-S-G-x-C.

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Protein Tyrosine Phosphatase (Y phosphatase). represents a polynucleotide encoding a protein tyrosine kinase. Tyrosine specific protein phosphatases (EC 3.1.3.48) (PTPase) (Fischer et al., Science (1991) 253:401; Charbonneau et al., Annu. Rev. Cell Biol. (1992) 8:463; Trowbridge, J. Biol. Chem. (1991) 266:23517; Tonks et al., Trends Biochem. Sci. (1989) 14:497; and Hunter, Cell (1989) 58:1013) catalyze the removal of a phosphate group attached to a tyrosine residue. These enzymes are very important in the control of cell growth, proliferation, differentiation and transformation. Multiple forms of PTPase have been characterized and can be classified into two categories: soluble PTPases and transmembrane receptor proteins that contain PTPase domain(s). Structurally, all known receptor PTPases are made up of a variable length extracellular domain, followed by a transmembrane region and a C-terminal catalytic cytoplasmic domain. PTPase domains consist of about 300 amino acids. The search of two conserved cysteines has been shown to be absolutely required for activity. Furthermore, a number of conserved residues in its immediate vicinity have also been shown to be important. The consensus pattern for PTPases is: [LIVMF]-H-C-x(2)-G-x(3)-[STC]-[STAGP]-x-[LIVMFY]; C is the active site residue. 35

Zinc Finger, C2H2 Type (Zincfing C2H2). SEQ ID NOs:308, 807, 1324, 1503, 1527, 3081, 3193 and 3306 correspond to polynucleotides encoding novel members of the of the C2H2 type zinc finger protein family. Zinc finger domains (Klug et al., Trends Biochem. Sci. (1987) 12:464; Evans et al., Cell (1988) 52:1; Payre et al., FEBS Lett. (1988) 234:245; Miller et al., EMBO J. (1985) 4:1609; and Berg, Proc. Natl. Acad. Sci. USA (1988) 85:99) are nucleic acid-binding protein structures. In addition to the conserved zinc ligand residues, it has been shown that a number of other positions are also important for the structural integrity of the C2H2 zinc fingers. (Rosenfeld et al., J. Biomol. Struct. Dyn. (1993) 11:557) The best conserved position is found four residues after the second cysteine; it is generally an aromatic or aliphatic residue. The consensus pattern for C2H2 zinc fingers is: C-x(2,4)-C-x(3)-[LIVMFYWC]-x(8)-H-x(3,5)-H. The two C's and two H's are zinc ligands.

Src homology 2. SEQ ID NOs:186, 2591, 3307 and 3339 represent polynucleotides encoding novel members of the family of Src homology 2 (SH2) proteins. The Src homology 2 (SH2) domain is a protein domain of about 100 amino acid residues first identified as a conserved sequence region between the oncoproteins Src and Fps (Sadowski I. et al., *Mol. Cell. Biol.* 6:4396-4408 (1986)). Similar sequences are found in many other intracellular signal-transducing proteins (Russel R.B. et al., *FEBS Lett.* 304:15-20 (1992)). SH2 domains function as regulatory modules of intracellular signalling cascades by interacting with high affinity to phosphotyrosine-containing target peptides in a sequence-specific and phosphorylation-dependent manner (Marangere L.E.M., Pawson T., *J. Cell Sci. Suppl.* 18:97-104 (1994); Pawson T., Schlessinger J., *Curr. Biol.* 3:434-442 (1993); Mayer B.J., Baltimore D., *Trends Cell. Biol.* 3:8-13 (1993); Pawson T., *Nature* 373:573-580 (1995)).

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The SH2 domain has a conserved 3D structure consisting of two alpha helices and six to seven beta-strands. The core of the domain is formed by a continuous beta-meander composed of two connected beta-sheets (Kuriyan J., Cowburn D., Curr. Opin. Struct. Biol. 3:828-837(1993)). The profile to detect SH2 domains is based on a structural alignment consisting of 8 gap-free blocks and 7 linker regions totaling 92 match positions.

Src homology 3. SEQ ID NO:234, 1832, and 1835 represent polynucleotides encoding novel members of the family of Src homology 3 (SH3) proteins. The Src homology 3 (SH3) domain is a small protein domain of about 60 amino acid residues first identified as a conserved sequence in the non-catalytic part of several cytoplasmic protein tyrosine kinases (e.g., Src, Abl, Lck) (Mayer B.J. et al.,

Nature 332:272-275 (1988)). Since then, it has been found in a great variety of other intracellular or membrane-associated proteins (Musacchio A. et al., FEBS Lett. 307:55-61 (1992); Pawson T., Schlessinger J., Curr. Biol. 3:434-442 (1993); Mayer B.J., Baltimore D., Trends Cell Biol. 3:8-13 (1993); Pawson T., Nature 373:573-580 (1995)).

The SH3 domain has a characteristic fold which consists of five or six beta strands arranged as two tightly packed anti-parallel beta sheets. The linker regions may contain short helices (Kuriyan J., Cowburn D., Curr. Opin. Struct. Biol. 3:828-837 (1993)).

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The function of the SH3 domain may be to mediate assembly of specific protein complexes via binding to proline-rich peptides (Morton C.J., Campbell I.D., Curr. Biol. 4:615-617 (1994)).

In general SH3 domains are found as single copies in a given protein, but there are a significant number of proteins with two SH3 domains and a few with 3 or 4 copies.

Fibronectin type III. SEQ ID NOs:746 and 1192 represent polynucleotides encoding novel members of the family of fibronectin type III proteins. A number of receptors for lymphokines, hematopoeitic growth factors and growth hormone-related molecules have been found to share a common binding domain. (Bazan J.F., Biochem. Biophys. Res. Commun. 164:788-795 (1989); Bazan J.F., Proc. Natl. Acad. Sci. U.S.A. 87:6934-6938 (1990); Cosman D. et al., Trends Biochem. Sci. 15:265-270 (1990); d'Andrea A.D., Fasman G.D., Lodish H.F., Cell 58:1023-1024 (1989); d'Andrea A.D., Fasman G.D., Lodish H.F., Curr. Opin. Cell Biol. 2:648-651 (1990)).

The conserved region constitutes all or part of the extracellular ligand-binding region and is about 200 amino acid residues long. In the N-terminal of this domain there are two pairs of cysteines known, in the growth hormone receptor, to be involved in disulfide bonds.

Two patterns detect this family of receptors. The first one is derived from the first N-terminal disulfide loop, the second is a tryptophan-rich pattern located at the C-terminal extremity of the extracellular region.

A consensus for this protein family is: C-[LVFYR]-x(7,8)-[STIVDN]-Cx-W (The two C's are linked by a disulfide bond]. A second consensus for this protein family is: [STGL]-x-W-[SG]-x-W-S.

LIM domain containing proteins. SEQ ID NOs:1269, 1309, 1360, and 1386 represent polynucleotides encoding novel members of the family of LIM domain containing proteins. A number of proteins contain a conserved cysteine-rich domain of about 60 amino-acid residues. (Freyd G. et al., Nature 344:876-879 (1990); Baltz R. et al., Plant Cell 4:1465-1466 (1992); Sanchez-Garcia I., Rabbitts T.H., Trends Genet. 10:315-320 (1994)).

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In the LIM domain, there are seven conserved cysteine residues and a histidine. The arrangement followed by these conserved residues is C-x(2)-C- x(16,23)-H-x(2)-[CH]-x(2)-C-x(2)-C-x(16,21)-C-x(2,3)-[CHD]. The LIM domain binds two zinc ions (Michelsen J.W. et al., Proc. Natl. Acad. Sci. U.S.A. 90:4404-4408 (1993)). LIM does not bind DNA, rather it seems to act as interface for protein-protein interaction. The consensus for this protein family is: C-x(2)-C-x(15,21)-[FYWH]-H-x(2)-[CH]-15 x(2)-C-x(2)-C-x(3)- [LIVMF]. The 5 C's and the H bind zinc.

C2 domain (protein kinase C like). SEQ ID NOs:1325 and 2282 represent polynucleotides encoding novel members of the family of C2 domain containing proteins. Some isozymes of protein kinase C (PKC) contain a domain, known as C2, of about 116 amino-acid residues, which is located between the two copies of the C1 domain (that bind phorbol esters and diacylglycerol) and the protein kinase catalytic domain. (Azzi A. et al., Eur. J. Biochem. 208:547-557 (1992); Stabel S., Semin. Cancer Biol. 5:277-284 (1994)).

The C2 domain is involved in calcium-dependent phospholipid binding (Davletov B.A., Suedhof T.C., J. Biol. Chem. 268:26386-26390 (1993)). domains related to the C2 domain are also found in proteins that do not bind calcium, other putative functions for the C2 domain include binding to inositol-1,3,5tetraphosphate. (Fukuda M., et al., J. Biol. Chem. 269:29206-29211 (1994).)

The consensus pattern for the C2 domain is located in a conserved part of that domain, the connecting loop between beta strands 2 and 3. The profile for the C2 domain covers the total domain. The consensus for this protein family is:: [ACG]-x(2)-L-x(2,3)-D-x(1,2)-[NGSTLIF]-[GTMR]-x-[STAP]-D-[PA]-[FY]

SEQ ID NO:1410 Serine proteases, trypsin family, active sites. represents a polynucleotide encoding a novel member of the family of serine protease, trypsin proteins. The catalytic activity of the serine proteases from the trypsin family is

provided by a charge relay system involving an aspartic acid residue hydrogen-bonded to a histidine, which itself is hydrogen-bonded to a serine. The sequences in the vicinity of the active site serine and histidine residues are well conserved in this family of proteases (Brenner S., *Nature 334*:528-530 (1988)).

A consensus for this protein family is: [LIVM]-[ST]-A-[STAG]-H-C [H is the active site residue]. A second consensus for this protein family is: [DNSTAGC]-[GSTAPIMVQH]-x(2)-G-[DE]-S-G-[GS]-[SAPHV]- [LIVMFYWH]- [LIVMFYSTANQH] [S is the active site residue].

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RNA Recognition Motif Domain (RRM, RBD, or RNP). SEQ ID NOs: 1464 and 1514 represent polynucleotides encoding novel members of the family of RNA recognition motif domain proteins (Bandziulis R.J. et al., Genes Dev. 3:431-437 (1989); Dreyfuss G. et al., Trends Biochem. Sci. 13:86-91 (1988)).

Inside the putative RNA-binding domain there are two regions which are highly conserved. The first one is a hydrophobic segment of six residues (which is called the RNP-2 motif); the second one is an octapeptide motif (which is called RNP-1 or RNP-CS). The position of both motifs in the domain is shown in the following schematic representation:

As a consensus pattern for this type of domain the RNP-1 motif was used. The consensus for this protein family is: [RK]-G-{EDRKHPCG}-[AGSCI]-[FY]-[LIVA]-x-[FYLM]

Phosphatidylinositol-specific phospholipase C, Y Domain. SEQ ID NO: 1707 represents a polynucleotide encoding a novel member of the phosphatidylinositol-specific phospholipase C, Y domain family of proteins. Phosphatidylinositol-specific phospholipase C (EC3.1.4.11), a eukaryotic intracellular enzyme, plays an important role in signal transduction processes (Meldrum E. et al., *Biochim. Biophys. Acta 1092*:49-71 (1991)). It catalyzes the hydrolysis of 1-phosphatidyl-D-myo-inositol-3,4,5- triphosphate into the second messenger molecules diacylglycerol and inositol-1,4,5-triphosphate. This catalytic process is tightly regulated by reversible phosphorylation and binding of regulatory proteins (Rhee S.G., Choi K.D., *Adv. Second Messenger Phosphoprotein Res. 26*:35-61 (1992); Rhee S.G., Choi K.D., *J. Biol. Chem. 267*:12393-12396 (1992); Sternweis P.C., Smrcka A.V., Trends Biochem. *Sci. 17*:502-506 (1992)).

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All eukaryotic PI-PLCs contain two regions of homology, referred to as "X-box" and "Y-box". The order of these two regions is the same (NH2-X-Y-COOH), but the spacing is variable. In most isoforms, the distance between these two regions is only 50-100 residues but in the gamma isoforms one PH domain, two SH2 domains, and one SH3 domain are inserted between the two PLC-specific domains. The two conserved regions have been shown to be important for the catalytic activity. At the C-terminal of the Y-box, there is a C2 domain possibly involved in Ca-dependent membrane attachment.

Serine Carboxypeptidases. SEQ ID NO:1744 represents a polynucleotide encoding a novel member of the serine carboxypeptidases family of proteins. Carboxypeptidases may be either metallo carboxypeptidases or serine carboxypeptidases (EC 3.4.16.5 and EC 3.4.16.6). The catalytic activity of the serine carboxypeptidases, like that of the trypsin family serine proteases, is provided by a charge relay system involving an aspartic acid residue hydrogen-bonded to a histidine, which is itself hydrogen-bonded to a serine (Liao D.I., Remington S.J., *J. Biol. Chem.* 265:6528-6531 (1990)).

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The sequences surrounding the active site serine and histidine residues are highly conserved in all these serine carboxypeptidases. A consensus for this protein family is: [LIVM]-x-[GTA]-E-S-Y-[AG]-[GS] [S is the active site residue]. A second consensus for this protein family is: [LIVF]-x(2)-[LIVSTA]-x-[IVPST]-x-[GSDNQL]-[SAGV]-[SG]-H-x- [IVAQ]-P-x(3)-[PSA] [H is the active site residue].

dsrm Double-Stranded RNA Binding Motif. SEQ ID NO:1818 represents a polynucleotide encoding a novel member of the dsrm double-stranded RNA binding motif proteins. In eukaryotic cells, a multitude of RNA-binding proteins play key roles in the posttranscriptional regulation of gene expression. Characterization of these proteins has led to the identification of several RNA-binding motifs. Several human and other vertebrate genetic disorders are caused by aberrant expression of RNA-binding proteins. (C. G. Burd & G. Dreyfuss, Science 265: 615-621 (1994)).

Proteins containing double stranded RNA binding motifs bind to specific RNA targets. Double stranded RNA binding motifs are exemplified by interferon-induced protein kinase in humans, which is part of the cellular response to dsRNA.

SEQ ID NOs:2577, 3183 and 3195 encode members of the 4 transmembrane integral membrane protein family. This family consists of type III proteins, which are integral membrane proteins that contain a N-terminal membrane-anchoring domain that is not cleaved during biosynthesis, and which functions as a translocation

signal and a membrane anchor. The proteins also have three additional transmembrane regions. The consensus pattern is: G-x(3)-[LIVMF]-x(2)-[GSA]-[LIVMF] (2)-G-C-x-[GA]-[STA]-x(20-[eG]-x(20-[CwN]-[LIVM](2).

SEQ ID NO:2944 encodes a polypeptide having a calpain large subunit, domain III. Calpains are a family of intracellular proteases that play a variety of biological roles. Calpain 3, also known as p94, is predominantly expressed in skeletal muscle and plays a role in limb-girdle muscular dystrophy type 2A. (Sorimachi, H. et al., Biochem. J. 328:721-732, 1997).

SEQ ID NOs:1911 and 1980 encode polypeptides having a C3HC4 type zinc finger domain (RING finger), which is a cysteine-rich domain of 40 to 60 residues that binds two atoms of zinc, and is believed to be involved in mediating protein-protein interactions. Mammalian proteins of this family include V(D)J recombination activating protein, which activates the rearrangement of immunoglobulin and T-cell receptor genes; breast cancer type 1 susceptibility protein (BRCA1); bmi-1 proto-oncogene; cbl proto-oncogene; and mel-18 protein, which is expressed in a variety of tumor cells and is a transcriptional repressor that recognizes and binds a specific DNA sequence. The consensus pattern is: C-x-H-x-[LIVMFY]-C-x(2)-C-[LIVMYA].

SEQ ID NO:3274 encodes a eukaryotic transcription factor with a fork head domain, of about 100 amino acid residues. Proteins of this group are transcription factors, including mammalian transcription factors HNF-3-alpha, -beta, and -gamma; interleukin-enhancer binding factor; and HTLF, which binds to a region of human T-cell leukemia virus long terminal repeat. The consensus pattern is [KR]-P-[PTQ]-[FYLVQH]-S-[FY]x(2)-[LIVM]-X(3,4)-[AC]-[LIM].

SEQ ID NO:3345 encodes a polypeptide having a PDZ domain. Several dozen signaling proteins belong to this group of proteins that have 80-100 residue repeats known as PDZ domains. Several of the proteins interact with the C-terminal tetrapeptide motifs X-Ser/Thr/X-Val-COO- of ion channels and/or receptors. (Ponting, C. P., Protein Sci. 6;464-468, 1997.)

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SEQ ID NO:3351 encodes a polypeptide in the family of phorbol esters/glycerol binding proteins. Phorbol esters (PE) are analogues of diacylglycerol (DAG) and potent tumor promoters. DAG activates a family of serine-threonine protein kinases, known as protein kinase C. The N-terminal region of protein kinase C binds PE and DAG, and contains one or two copies of a cysteine-rich domain of about 50 amino acid residues. Other proteins having this domain include diacylglycerol kinase; the vav oncogene; and N-chimaerin, a brain-specific protein. The DAG/PE binding

domain binds two zinc ions through the six cysteines and two histidines that are conserved in the domain. The consensus pattern is: H-x-[LIVMFYW]-x(8, 11)-C-x(2)-C-x-(3)-[LIVMFC]-x(5, 10)-C-x(2)-C-x(4)-[HD]-x(2)-C-x(5, 9)-C.

SEQ ID NO:2216 encodes a polypeptide having a WW/rsp5/WWP domain. The protein is named for the presence of conserved aromatic positions, generally tryptophan, as well as a conserved proline. Proteins having the domain include dystrophin, vertebrate YAP protein, and IQGAP, a human GTPase activating protein which acts on ras. The consensus pattern is: W-x(9,11)-[VFY]-[FYW]-x(6,7)-[GSTNE]-[GSTQCR]-[FYW]-x(2)-P.

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SEQ ID NO:2428 encodes a member of the dual specificity phosphatase family, having a catalytic domain, and SEQ IDS NOs:2281 and 2310 encode members of the protein tyrosine phosphatase family. These families are related and classified as tyrosine specific protein phosphatases. The enzymes catalyze the removal of a phosphate group from a tyrosine residue, and are important in the control of cell growth, proliferation, differentiation, and transformation. The consensus pattern is [LIVMF]-H-C-x(2)-G-x-(3)-[STC]-[STAGP]-x-[LIVMFY].

Table 1

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3	427782	RTA00002686F1.00.11F.Seq RTA00002712F.a.06.1.P.Seq	F	M00023282A:C02	CH04MAL
4	29372	RTA00002712F.3.00.1.F.3eq RTA00002694F.b.02.1.P.Seq	F	M00043419D:A10	CH20COHLV
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9	20652	RTA00002710F.k.01.1.P.Seq RTA00002663F.k.18.!.P.Seq	F	M00022767B:G11	CH03MAH
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15	46779	RTA00002711F.c.14.1.P.Seq		M00039909C:G05	CH09LNL
16	375772	RTA00002681F.p.01.2.P.Seq		M00033243B:A05	CH08LNH
17	430689	RTA00002669F.j.01.3.P.Seq		M00039345C:C12	CH09LNL
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59	375666	RTA00002677F.n.16.2.P.Seq	F	M00027292D:F10	CH04MAL
60	162369	RTA00002713F.d.24.1.P.Seq	F	M00005531D:F06	CH02COH
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72	449831	RTA00002691F.a.17.3.P.Sec	F	M00039274B:G07	CH09LNL
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74	20599	RTA00002708F.h.06.1.P.Sec	F F	M00037652B:F11	CH0+MAL
75	41115	RTA00002713F.o.11.1.P.Sec	F	M00004278A:F09	CH01COH
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84		RTA00002673F.n.11.1.P.Se	a F	M00039108D:B06	CH09LNL
85		RTA00002683F.c.09.1.P.Se	q F	M00040039D:D06	CH08LNH
86		RTA00002669F.h.18.2.P.Se	q F	M00033226A:A11	CH09LNL
87		RTA00002674F.1.17.1.P.Se	q F	M00039166B:G06	CH09FNF
88		RTA00002672F.p.20.1.P.Se	eq F	M00039049D:G07	CH09LNL
89		RTA00002672F.p.22.1.P.Se	eq F	M00039050A:H10	CH09LNL
90		RTA00002678F.h.17.2.P.Se	q F	M00039476B:A02	CH09FNF
—		RTA00002672F.b.20.1.P.Se	q F	M00038638D:H03	
9		RTA00002676F.p.07.1.P.Se	q F	M00039328D:D07	CHOOLNL
9:		RTA00002672F.c.08.1.P.Se	q F	M00038661A:A07	CH09LNL
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96	375339	RTA00002678F.m.23.2.P.Seq	F	M00039616A:B10	CH03MAH
97	14197	RTA00002710F.f.15.1.P.Seq	F	M00022084D:B01	CH09LNL
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	68 373282	BL700005080F1912		M00004168A:G1	CHOICOR
	69 33397	B T 4 0000 266 11. 11. 104. 1. 1	1011	M00039213B:F05	CHO9LNL
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407		RTA00002714F.c.10.1.P.Se	q F	M00027836D:F12	CH08LNH
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41		RTA00002669F.k.08.2.P.S	eq F	M00033263B:G04	
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46	6 431370	RTA00002669F.m.04.2.P.5		M0001-4-6.4:C09	CH0+W-71
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47		RTA00002713F.i.20.1.P.S	sed L	144002	<u>.</u>

					
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473	379684	RTA00002691F.f.12.2.P.Seq	F	M00043411D:H06	CH17COHLV
474	451564	RTA00002710F.a.15.1.P.Seq	F	M00007943D:C09	CH03MAH
475	7571	RTA00002713F.k.21.1.P.Seq	F	M00027525B:D06	CH04MAL
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477	12960	RTA00002713F.o.05.1.P.Seq	F	M00027641C:A03	CH04MAL
478	186730	RTA00002713F.m.24.1.P.Seq		M00027596C:E06	CH04MAL
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489	447544	RTA00002686F.c.23.1.P.Seq		M00040141 D:F05	CHISEDT
490	401872	RTA00002686F.C.25.111.Seq RTA00002674F.g.19.2.P.Seq		M00039139A:C09	CH09LNL
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510		RTA00002710F.p.05.1.P.Set		M00021972D:C11	СНОЗМАН
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55		RTA00002674F.m.12.1.P.S		M00039066D:G08	CH09LNL
55		RTA00002673F.d.24.2.P.S		M00039066D:G08	CH09LNL
55		RTA00002673F.e.01.2.P.S	eq F	M00039686C:E06	CH09LNL
55		RTA00002679F.i.03.1.P.S	eq F	M00039792A B04	CH09LNL
56		RTA00002680F.e.15.1.P.S	eq F	M00034792A:B04	CH03MAH
56		RTA00002710F.n.23.1.P.S	eq F	MO0022071B400	CH03MAH
		RTA00002663F.n.15.1.P.S	eq t	M100023039D:305	CHOSLNH
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567	379207	RTA00002665F.k.19.1.P.Seq	F	M00031419D:C04	CH08LNH
568	427893	RTA00002684F.g.19.2.P.Seq	F	M00040305.A:D11	CH09LNL
569	377530	RTA00002668F.c.11.1.P.Seq	F	M00032918C:B10	CH08FNH
570	429707	RTA00002665F.i.04.1.P.Seq	F	M00028770A:D04	CH08LNH
571	427610	RTA00002662F.b.22.2.P.Seq	F	M00006680B:D02	CH02COH
572	100699	RTA00002602F.0.22.2.1.1.P.Seq		M00040017A:C06	CH09LNL
573	378974	RTA00002682F.m.21.1.P.Seq		M00039127D:E10	CH09LNL
574	373607	RTA00002674F.d.15.1.P.Seq		M00028215D:F03	CH08LNH
575	262951	RTA00002665F.d.04.3.P.Seq		M00027301B:B08	CH04MAL
576	30748	RTA00002713F.e.11.1.P.Seq		M00027837C:D09	CH04MAL
577	161116	RTA00002714F.c.11.1.P.Seq		M00040029.A:G04	CH09LNL
578	379211	RTA00002682F.p.20.1.P.Seq		M00033243B:A05	CH08LNH
579	430689	RTA00002669F.i.24.1.P.Seq		M00039104D:C09	CH09LNL
580	374122	RTA00002673F.1.22.2.P.Seq	F	M00039398A:B10	CH09LNL
581	376521	RTA00002677F.h.06.2.P.Sec	F	M00033308B:G05	CH09LNL
582	372834	RTA00002670F.b.12.2.P.Sec	F	M00040022C:D06	CH09LNL
583	379014	RTA00002682F.o.02.1.P.Sec	1 F	M00039340B:G08	CH09LNL
584	376344	RTA00002677F.b.18.2.P.Sec	1 F		CH09LNL
	376485	RTA00002676F.f.01.2.P.Sec	1 F	M00039288C:B11	CH02COH
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588		RTA00002714F.c.20.1.P.Se	q F	M00028120D:F12	CH04MAL
589		RTA00002713F.c.20.1.P.Se	q F	M00027263A:F10	CH04MAL
590		RTA00002670F.i.22.2.P.Se	q F	M00033432B:H10	
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592		RTA00002663F.f.10.1.P.Se	a F	M00022233C:A12	CH03MAH
593		RTA00002671F.m.02.1.P.Se	7	M00038328D:A03	CH09LNL
594		RTA00002712F.k.23.1.P.Se		M00027022D:G11	CH04MAL
595	26429	RTA00002712F.K.25.1.1.5		M00022979.A:D05	CH03MAH
596	17983	RTA00002711F.f.10.1.P.Se	7	M00039888B:D03	CH09LNL
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598	63005	RTA00002712F.m.21.1.P.S	77	M00005384.A:C11	CH02COH
599	23030	RTA00002709F.b.10.2.P.S	*71	M00033457D:A05	CH09LNL
600	0 372946	RTA00002670F.1.07.2.P.Se		M00039792A:B04	CH09LNL
60		RTA00002680F.e.15.2.P.S		M00039080C:H06	CH09LNL
60		RTA00002673F.i.08.1.P.Se	eq F	M00039938C:A08	CH09LNL
60		RTA00002682F.e.09.1.P.S	eq F	M00039257D:C03	CH09LNL
60		RTA00002675F.n.15.1.P.S	eq F	M00039237D:C03	CH09LNL
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613	46804	RTA00002712F.i.18.1.P.Seq	F	M00026935C:B04	CH04MAL
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615	375285	RTA00002670F.j.13.2.P.Seq	F	M00033437C:C03	CH09LNL
616	373000	RTA00002670F.J.15.2.1 .5eq	F	M00039869B:F06	CH09LNL
617	378679	R [A00002081F.1.10.1.1.5eq	F	M00027016A:B06	CH04MAL
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620	186425	RTA00002713F.c.04.1.P.Seq	F	M00039288C:B11	CH09LNL
621	376485	RTA00002676F.e.24.2.P.Seq		M00027108C:B03	CH04MAL
622	41108	RTA00002712F.n.12.1.P.Seq	F	M00033186C:D11	CH08LNH
623	430876	RTA00002669F.c.02.1.P.Seq	F	M00027537C:B01	CH04MAL
624	185716	RTA00002713F.1.07.1.P.Seq		M00023333D:C12	CH04MAL
625	85338	RTA00002712F.b.18.1.P.Seq		M00027596A:A10	CH04MAL
626	185597	RTA00002713F.m.23.1.P.Seq		M00027526D:F03	CH04MAL
627	139348	RTA00002713F.k.23.1.P.Seq		M00043164C:E12	CH19COP
628	454665	RTA00002693F.d.15.2.P.Seq	F	M00027528C:B10	CH04MAL
629	186387	RTA00002713F.I.01.1.P.Seq	F	M00027528C:B10	CH04MAL
630	186387	RTA00002713F.k.24.1.P.Seq	F	M00004308C:C06	CH01COH
631	21093	RTA00002708F.h.20.1.P.Sed	F	M00004308C:C00	СНОЗМАН
632	20827	RTA00002710F.c.23.1.P.Sec	F	M00021871D:112	CH04MAL
633	21290	RTA00002712F.f.24.1.P.Seq	F		СНОЗМАН
634	17646	RTA00002710F.d.22.1.P.Sec	F	M00021908D:G12	CHISEDT
635	402817	RTA00002686F.a.10.1.P.Sec	I F	M00039736D:G08.	CH04MAL
636		RTA00002713F.n.09.1.P.Sec	ı F	M00027615A:F10	CH04MINE
637		RTA00002669F.c.02.3.P.Sec	1 <u>F</u>	M00033186C:D11	CH09LNL
638		RTA00002679F.a.21.2.P.Sec	F F	M00039655C:E08	CH09LNL
639		RTA00002674F.m.03.2.P.Se	q F	M100039168C:A04	CHOICOH
		RTA00002708F.i.06.1.P.Sec	al <u>F</u>	M00004340C:C07	CH20COHLV
640		RTA00002694F.d.05.1.P.Se	q F	M00043492A:E01	
641		RTA00002672F.i.05.2.P.Set	q F	M00039014A:H10	CH09LNL
642		RTA00002683F.o.21.2.P.Se	q F	M00040100D:B06	CH09LNL
643		RTA00002671F.m.02.2.P.Se	g F	. M00038328D:A03	CH09LNL
644		RTA00002681F.m.20.1.P.Se	a F	M00039898.4:A08	CH09LNL
645		RTA00002669F.f.07.2.P.Se	a F	M00033204B:A07	CH08LNH
646		RTA00002683F.a.06.1.P.Se	1	M00040032A:B03	CH09LNL
647		RTA00002683F.a.00.111.30	7]	M00001439C:H06	CH01COH
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649	374410	RTA00002674F.k.11.1.P.Se		M00039118D:A06	CH09LNL
650		RTA00002674F.a.01.2.P.Se		M00033202D:G06	CH08LNH
65		RTA00002669F.e.23.2.P.Sc		M00038995 D: E05	CH09LNL
65	2 240615	RTA00002672F.e.19.1.P.Se		M00032545B:H09	CH08LNH
65		RTA00002666F.d.01.1.P.S		M00032545 D:G05	CH08LNH
65		RT.A00002666F.d.03.1.P.S	eq F	M00021955A:H02	CH03MAH
65		RTA00002710F.e.11.1.P.S	eq F	M00022175D:D12	CH03MAH
65		RTA00002710F.g.11.1.P.S	eq F	M0002218-C:C11	CH03M.AH
65		RTA00002710F.g.18.1.P.S	eq F	M00039875D:A10	CH09LNL
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SEQ	CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
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660	15596	RTA000027101.g.	F	M00007108B:A02	CH02COH
661	21028	RTA00002672F.h.15.2.P.Seq	F	M00039011D:C10	CH09LNL
6 62	374063	RTA00002684F.a.03.2.P.Seq	F	M00040107B:H07	CH09LNL
663	380686	RTA00002686F.g.11.1.P.Seq	F	M00040181B:H09	CHISEDT
664	402950	RTA0000_686F.L04 LP Seq	F	M00031485D:G02	CH08LNH
665	428064	RTA00002665F.1.04.1.P.Seq	F	M00004046C:A08	CH01COH
666	23310	RTA00002708F.e.10.1.P.Seq	F	M00039339C:F03	CH09LNL
667	376233	RTA00002677F.b.15.2.P.Seq	F	M00039168C:A04	CH09LNL
668	375848	RTA00002674F.m.03.1.P.Seq	F	M00028772C:B09	CH08LNH
669	242251	RTA00002665F.i.08.1.P.Seq	F	M00038999D:C11	CH09LNL
670	374064	RTA00002672F.f.15.2.P.Seq	F	M00022099B:D06	CH03MAH
671	146260	RTA00002663F.d.17.1.P.Seq	F	M00039385B:E09	CH09LNL
672	375575	RTA00002677F.e.22.2.P.Seq		M00028201B:H12	CH08LNH
673	355518	RTA00002665F.c.15.3.P.Seq		M00005539D:G01	CH02COH
674	184223	RTA00002662F.b.08.2.P.Seq	F	M00027078A:B02	CH04MAL
675	213306	RTA00002664F.e.07.2.P.Seq	F	M00032907 A:G04	CH08LNH
676	429566	RTA00002668F.b.04.1.P.Seq	F	M000329074:F04	CH09LNL
677	378656	RTA00002682F.c.09.1.P.Seq	F	M000379277415 M00032940A:C02	CH08LNH
678	427760	RTA00002668F.e.23.1.P.Seq	<u> </u>	M00040032A:B03	CH09LNL
679	372795	RTA00002683F.a.06.2.P.Seq	F	M00032577A:C04	CH08LNH
680	429340	RTA00002666F.f.12.1.P.Seq	F-	M00032977A:C07	CH08LNH
681	429822	RTA00002668F e.17.1.P.Seq	l <u> </u>	M00032737B:207	CH09LNL
682	375224	RTA00002680F.d.22.2.P.Sec	F	M00039733B::100	CH09LNL
683	378347	RTA00002681F.h.07.1.P.Sec	1 - F	M00039873B:R13	CH09LNL
684	380109	RTA00002682F.i.17.1.P.Sec	F	M00040097A:C12	CH09LNL
685	379001	RTA00002683F.o.02.1.P.Sec	9 F	M0004004;A:C12	CH09LNL
686	375348	RTA00002676F.i.12.3.P.Sec	1 F	M00039304B:B07	CH09LNL
	377889	RTA00002672F.c.08.2.P.Se	q F	M00038661.4407	CH08LNH
687		RTA00002667F.g.05.1.P.Se	9 <u>F</u>	M00032793A:F06	CH09LNL
688		RTA00002682F.I.24.1.P.Set	q <u>F</u>	M00040014B:D01	CH09LNL
689		RTA00002681F.m.22.2.P.Se	q F	M00039898D:C06	CH03MAH
690		RTA00002710F.j.21.1.P.Se	9 -	M00022433A:E02	CH09LNL
691		RTA00002680F.f.03.1.P.Se	q F	M100039793 D:C05	CH09LNL
692		RTA00002681F.m.20.2.P.Sc	eq F	M00039898.A:A08	CH08LNH
693	170410	RTA00002667F.e.09.1.P.Se	q F	M00032766C:A04	CH03MAH
694		RTA00002710F.i.10.1.P.Se	q F	M00022363C:G12	CH04MAL
69:	27.60	RTA00002713F.e.23.1.P.Sc	eq F	M00027314C:D09	CH04MAE
690		RTA000027757	eq F	M100033140D:F06	
69		RTA00002670F.h.23.2.P.S	eq F	M00033424B:A04	CH09LNL
69		RTA00002709F.f.07.1.P.S	eq F	M00006549D:B02	CH02COH
69		RTA00002707Fit.03.2.P.S	ea F	M00033424D:H12	CH09LNL
70		RTA00002671F.d.20.1.P.S		M00038272D:F11	CH09LNL
70		RTA00002684F.h.06.2.P.S	- 3	M00040307B:F01	CH09LNL
70			~ ~	M00040302C::A04	CH09LNL
70		- 1 · · · · · · · · · · · · · · · ·		M00039286A:C06	CH09LNL
70	374747	RT.A00002676F.e.07.2.P.S		M00027080A:B01	CH04MAL
70	5 185848	RTA00002712F.m.11.1.P.5	<u> </u>		•

SEQ			COLENIA TON	CLONE ID	LIBRARY
ID	CLUSTER	320	ORIENTATION	M00039287C:A06	CH09LNL
706	374311	RTA00002676F.e.18.2.P.Seq	F	M00032726C:C01	CH08LNH
707	278923	RTA00002667F.b.10.1.P.Seq	F	M00032726C:C01	CH09LNL
708	378667	RTA00002681F.b.11.2.P.Seq	F	M00039084D:D07	CH09LNL
709	380454	RTA00002673F.j.16.1.P.Seq	F	M00039084D:D07 M00033425A:C10	CH09LNL
710	381576	RTA00002670F.i.04.2.P.Seq	F	M00039260C:G03	CH09LNL
711	375067	RTA00002675F.o.03.1.P.Seq	F	M00039260C:G03 M00027741B:F09	CH04MAL
712	89706	RTA00002714F.a.11.1.P.Seq	F	M000277415:107	CH03MAH
713	10583	RTA00002711F.h.11.1.P.Seq	F	M00023100A:E12	CH09LNL
714	379982	RTA00002682F.i.16.1.P.Seq	F	M00039987C:212 M00039828B:C05	CH09LNL
715	378532	RTA00002680F.n.04.3.P.Seq	F	M00039774C:A03	CH09LNL
716	379776	RTA00002680F.a.22.2.P.Seq	F	M00039774C::403	CH09LNL
717	374136	RTA00002673F.f.16.1.P.Seq	F	M00039072C:C03	CH03MAH
718	98471	RTA00002663F.j.21.1.P.Seq	F	M00022870B:E10	CH08LNH
719	125365	RTA00002668F.j.07.1.P.Seq	F	M00033019B.E10	CH09LNL
	375431	RTA00002680F.f.03.2.P.Seq	F	M00039793D:C05	CHOICOH
720	62826	RTA00002661F.g.20.1.P.Seq	F	M00004105D:D05	CH09LNL
721	379972	RTA00002679F.e.10.1.P.Seq	F	M00039672D:D10	CH09LNL
722	377554	RTA00002679F.f.10.1.P.Seq	F	M00039675D:B03	CH04MAL
723	230479	RTA00002664F.c.16.2.P.Sec	F	M00026915B:C06	СНОЗМАН
724		RTA00002663F.j.19.1.P.Seq	F	M00022668B:B12	CH09LNL
725		RTA00002679F.h.18.1.P.Sec	F	M00039684D:B08	CH09LNL
726		RTA00002679F.a.10.2.P.Sec	F	M00039652B:D05	CH03MAH
727		RTA00002663F.j.07.1.P.Sec	F	M00022640C:C12	CH14EDT
728		RTA00002687F.d.01.2.P.Sec	q <u>F</u>	M00039945C:F09	CH12EDT
729		RTA00002685F.b.18.2.P.Se	q F	M00039371B:H06	CH03MAH
730		RT.A00002663F.k.10.1.P.Se	q F	M00022731A:D02	CH04MAL
731		RTA00002713F.c.18.1.P.Se	q F	M00027258.A::A07	CHUANAL
732		RTA00002686F.b.09.1.P.Se	q F	M00039756B:H06	CH20COHLV
733		RTA00002694F.e.06.1.P.Se	q F	M00043634A:C10	CH04MAL
73-		RTA00002713F.c.16.1.P.Se	g F	M00027256B:H09	
73:		RTA000027131.a.15.2.P.Se	g F	M00042626B:D08	CH18CON
730		RTA00002682F.e.23.1.P.Sc	g F	M00039940D:G08	CH09LNL
73		RTA00002680F.n.04.2.P.Sc	g F	M00039828B:C05	CH09LNL
73		RTA00002713F.b.22.1.P.S	eq F	M00027233B:C01	CH04MAL
73		RTA00002713F.0.22.11.	F F	M00027179D:E06	CH04MAL
74	0 9332	RTA00002712F.p.10.11.3 RTA00002687F.d.04.2.P.S		M00039947.A:D06	CH14EDT
74	1 240318	RTA00002687F.d.04.2.1.3 RTA00002687F.c.11.2.P.S	ea F	M00039942D:C01	CH14EDT
74		RT.A00002687F.C.11.2.1.5		M00026868C:E11	CH04MAL
74		RTA00002712F.g.09.1.P.S		M00026856D:F02	CH04MAL
74	14 185642	RTA00002712F.f.20.1.P.S		M00042905D:D02	CHISCON
70	15 447544	RTA00002689F.e.18.3.P.S		M00039766.A:G07	CH14EDT.
74	46 403274	RTA00002687F.b.10.2.P.S		M00040208A:C03	CHIJEDT
74	47 404257	RTA00002687F.g.06.1.P.S	7 7 1	M00040318C:H11	CH14EDT
-	43 403868	RTA00002687F.k.05.2.P.S	***	M00043392D:C11	CH17COHL
	49 450074	RTA00002691F.e.12.2.P.S		M100040202A:F05	CH14EDT
-	50 404520	RTA00002687F.f.05.2.P.S	eq F	M00042956C:B06	CHISCON
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	52 455178	RTA00002694F.b.19.1.P.	Seq F	W100042447.A.CO.	

SEQ		aro vivir	ORIENTATION	CLONE ID	LIBRARY
ID	CLUSTER	300	F	M00042595A:B01	CH20COHLV
753	455136	RTA00002694F.a.08.1.P.Seq		M00040097.A:C12	CH09LNL
754	379001	RTA00002683F.o.02.2.P.Seq	F	M00039118B:C05	CH09LNL
755	374763	RTA00002673F.p.21.1.P.Seq		M00040281D:B01	CHISEDT
756	402508	RTA00002686F.o.15.1.P.Seq	F	M00033288B:D12	CH08LNH
757	431370	RTA00002669F.m.04.3.P.Seq	F	M00033583B:E06	CH09LNL
758	380500	RTA00002670F.p.19.1.P.Seq	F	M00039461A:F04	CH09LNL
759	376743	RTA00002678F.e.22.2.P.Seq	F	M00039107C:E04	CH09LNL
760	191690	RTA00002673F.m.19.1.P.Seq	F	M000391070:E09	CH09LNL
761	374264	RTA00002671F.p.21.2.P.Seq	F	M00038020D:209 M00033595A:C11	CH09LNL
762	373020	RTA00002671F.b.20.2.P.Seq	F	M00033393A.C11	CH09LNL
	375231	RTA00002671F.m.20.2.P.Seq	F.	M00038387B:A07	CH02COH
763	16180	RTA00002709F.j.17.1.P.Seq	<u> </u>	M00006977D::A03	CH09LNL
764		RTA00002683F.c.17.2.P.Seq	F	M00040041C:C09	CH09LNL
765	379403	RTA00002677F.d.24.2.P.Seq	F	M00039381D:C02	CH09LNL
766	375382	RTA00002683F.c.03.2.P.Seq	F	M00040038D:G04	CH09LNL
767	379653	RTA00002681F.e.14.2.P.Seq	F	M00039864A:A07	CHOSLNH
768	377858	RTA00002668F.h.18.1.P.Seq	F	M00032995C:C05	
769	430861	RTA00002677F.a.11.2.P.Seq	F	M00039334B:E03	CH09LNL
770	376128	RTA00002676F.n.20.2.P.Sec	F	M00039322A:F04	CH09LNL
771	375009	RTA00002667F.n.22.1.P.Sec	1	M00032871D:E11	CH08LNH
772	429816	RTA00002607F.II.22.1.1.566		M00039877C:C03	CH09LNL
773	375657	RTA00002681F.h.13.2.P.Sec	1	M00032530D:C02	CH08LNH
774	427889	RTA00002666F.b.14.1.P.Sec	<u> </u>	M00039391D:F08	CH09LNL
775	376761	RTA00002677F.g.03.2.P.Se	<u> </u>	M100040115B:A04	CH09LNL
776	44025	RTA00002684F.b.24.2.P.Se		M100040115B:A04	CH09LNL
777		RTA00002684F.c.01.2.P.Se		M00039909D:C02	CH09LNL
778		RTA00002681F.p.04.2.P.Se	71	M00028185B:A06	CH08LNH
779		RTA00002665F.b.13.1.P.Se	3	M00039068C:E06	CH09LNL
780		RTA00002673F.e.12.1.P.Se		M00039797C:G05	CH09LNL
781		RTA00002680F.g.09.1.P.Se	71	M00027299B:B12	CH04MAL
782		RTA00002664F.g.08.2.P.Se	q F	M00039339.4:H07	CH09LNL
783		RTA00002677F.b.14.2.P.Se	q F	M00039407B:G02	CH09LNL
78-		RTA00002677F.j.11.2.P.Se	<u>q </u>	M00039384C:E02	CH09LNL
78:		RTA00002677F.e.16.2.P.Se	eq F	M00005445D:B01	
78		RTA00002662F.b.01.2.P.S	eq F	M000034430:B01	
		RTA00002673F.e.12.2.P.S	eq F		
78	270051	RTA00002667F.b.23.1.P.S	eq F	M00032731B:C10	
78		RTA00002682F.m.19.1.P.S	[P96]	M00040016C:H12	
78		RTA00002710F.1.05.1.P.S	eq F	M00022533A:A08	
79		RTA00002664F.e.23.2.P.S	eq F	M00027162B:F05	
79		RTA00002683F.c.20.2.P.S	eq F	M00040042B:A10	
79		RTA00002665F.b.13.3.P.S	eq F	M00028185B:A00	
79		RTA00002673F.j.12.2.P.S	eq F	M00039084C:G0	7 CH09LNL
79	380618	RTA00002667F.g.16.1.P.S		M00032797B:G03	2 CH08LNH
79	35646	R1A0000_00/F.g.10.1.F.		M00028196D:A0	3 CHOSLNH
79	96 46407	RTA00002665F.c.10.3.P.S		M00039124C:F0	CHOOLNL
7	97 373720		²⁶ 91	M00032944B:B0	2 CHOSLNH
F-7	98 429693	RTA00002668F.f.05.1.P.: RTA00002678F.p.04.2.P.		M00039636C:D1	I CHOOLNL

SEQ		SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
[D]	CLUSTER	RTA00002681F.h.13.1.P.Seq	F	M00039877C:C03	CH09LNL
800	375657	RTA00002681F.n.13.1.1.5eq	F	M00039063B:D08	CH09LNL
801	374868	RTA00002673F.d.08.2.P.Seq	F	M00032766B:D12	CH08LNH
802	428716	RTA00002667F.e.08.1.P.Seq	F	M00040115B::A04	CH09LNL
803	44025	RTA00002684F.c.01.1.P.Seq	F	M00033033C:H01	CH08LNH
804	430327	RTA00002668F.k.11.1.P.Seq	F	M00039061B:F08	CH09LNL
805	374328	RTA00002673F.c.24.1.P.Seq	F	M00040019A:E01	CH09LNL
806	376946	RTA00002682F.n.10.1.P.Seq	F	M00039420D:D03	CH09LNL
807	375522	RTA00002677F.n.08.2.P.Seq		M00039767B:A04	CH14EDT
808.	395617	RTA00002687F.b.15.2.P.Seq	F F	M000377672::A06	CH04MAL
809	21686	RTA00002712F.g.05.1.P.Seq		M00042623D:D07	CH18CON
810	452038	RTA00002692F.a.09.2.P.Seq	F	M00042823B:B07 M00023042D:D02	CH03MAH
811	25632	RTA00002711F.g.16.1.P.Seq	F	M00023042D:D02	CH03MAH
812	152487	RTA00002663F.e. 12.1.P.Seq	- F	M00022181C.D01	CH09LNL
813	378226	RTA00002680F.g.09.2.P.Seq	F	M00039797C:G05	CHISEDT
814	402446	RTA00002686F.c.04.1.P.Seq	<u> </u>	M00040133B:B03	CH14EDT
	403642	RTA00002687F.c.24.2.P.Seq	<u>F</u>	M00039945C:F09	CH04MAL
815	186359	RTA00002713F.g.24.1.P.Seq	F	M00027379C:B07	CH14EDT
816	404290	RTA00002688F.e.04.2.P.Sec	F	M00040395B:D11	CH09LNL
817		RTA00002676F.g.19.2.P.Sec	F	M00039298B:D03	CH09LNL
818	375443 380279	RTA00002673F.i.24.1.P.Sec	F	M00039082B:A05	CHI4EDT
819		RTA00002687F.e.06.1.P.Sec	F	M00039955C:C04	
820	386110	RTA00002673F.j.01.1.P.Sec	F	M00039082B:A05	CH09LNL
821	380279	RTA00002675F.p.06.1.P.Se	F	M00039266.4:B02	CH09LNL
822	386986	RTA00002713F.h.01.1.P.Se	g F	M00027379C:B07	CH04MAL
823		RTA000027751 RTA00002677F.o.20.2.P.Se	a F	M00039425D:E12	CH09LNL
824		RTA00002679F.h.01.1.P.Se	a F	M00039681B:H09	CH09LNL
825		RTA00002684F.b.24.1.P.Se	g F	M00040115B::A04	CH09LNL
826		RTA00002711F.c.12.1.P.Se	a F	M00022854A:B03	CH03MAH
827		RTA000027111.c.12.43.66 RTA00002687F.g.03.2.P.Se	7	M00040207B:D08	CH14EDT
828		RTA00002687F.f.07.1.P.Se		M00040203A:H06	CH14EDT
829		RTA00002693F.b.12.2.P.Se	7	M00043093C:G11	CH19COP
830	454806	RTA00002674F.f.21.2.P.Se		M00039135D:G02	CH09LNL
831	376829	RTA00002674F.1.21.2.1.3c		M00043518B:D06	CH20COHLV
832	456309	RTA00002694F.d.16.1.P.Sc	7	M00039015D:H04	CH09LNL
83:	374510	RTA00002672F.i.17.2.P.Se	7	M00040090B:G09	CH09LNL
834	377232	RTA00002683F.m.08.2.P.S	731	M00039025A:H09	CH09LNL
83	5 375779	RTA00002672F.j.20.2.P.Sc		M00033585D:A02	CH09LNL
83	6 90746	RTA00002671F.a.07.2.P.S		M00042970C:H10	CH18CON
83	7 453002	RTA00002692F.b.21.2.P.S		M00040273B:H12	CHISEDT
83		RTA00002686F.n.12.1.P.S		M00040286C:C02	CH13EDT
83		RTA00002686F.p.07.1.P.S	eq F	M00039533D:F04	CHIZEDT
84		RTA00002685F.i.07.1.P.S	eq F	M00040233C:G05	CHISEDT
84		RTA00002686F.j.18.1.P.S	eq F	M100033427D:F01	CH09LNL
84		RTA00002670F.i.16.1.P.S	eq F	M00039425C:G01	CH09LNL
84		RTA00002677F.o.18.2.P.S	seq F	M00039423C.G01	CH08LNH
84		RTA00002667F.c.01.1.P.S	seq F	M00032731C:C07	CH09LNL
8-		RTA00002675F.h.01.1.P.S	Seq F	M00039230D:D09	CH03MAH
<u> </u>	6 20162	RTA00002710F.n.20.1.P.5	Seq F	M00022662D:G11	<u> </u>

SEQ		570 314315	ORIENTATION	CLONE ID	LIBRARY
ID]	CLUSTER	360	F.	M00039381C:H08	CH09LNL
847	375782	RTA00002677F.d.23.2.P.Seq	F	M00038639D:F07	CH09LNL
343	372958	RTA00002672F.c.02.1.P.Seq	F	M0004038 D:H05	CHIJEDT
849	403940	RTA00002688F.d.07.2.P.Seq	F	M00023020C:G08	CH03MAH
850	8490	RTA00002711F.g.03.1.P.Seq		M00039230 D: D09	CH09LNL
851	374809	RTA00002675F.g.24.1.P.Seq	F	M00040305C:H06	CH09LNL
852	377788	RTA00002684F.g.24.2.P.Seq	F	M00022976C:F04	СНОЗМАН
853	13847	RTA00002711F.f.09.1.P.Seq	F	M00022978C:104	CH09LNL
854	374172	RTA00002673F.k.16.1.P.Seq	F		CH09LNL
855	380314	RTA00002682F.1.07.1.P.Seq	F	M00040009D:B07	CH04MAL
856	47231	RTA00002714F.b.15.1.P.Seq	F	M00027813C:F01	CH12EDT
857	400287	RTA00002685F.k.10.1.P.Seq	F	M00039584C:C01	CHIZEDT
	400533	RTA00002685F.a.02.2.P.Seq	F	M00039181D:E05	CH15CON
858		RTA00002689F.c.07.1.P.Seq	F	·M00042696B:E05	CH03MAH
859	447594· 147357	RTA00002711F.e.15.1.P.Seq	F	M00022928B:C01	
860		RTA00002685F.o.22:2.P.Seq	F	M00039642D:B12	CHIZEDT
861	401141	RTA00002687F.c.03.1.P.Seq	F	M00039770A:G11	CH14EDT
862	404620	RTA00002709F.1.20.1.P.Seq	F	M00007149.A:G02	CH02COH
863	24360	RTA000027071112011111111111111111111111111	F	M0003908-C:G07	CH09LNL
864	380618	RTA000026791 J. 12-13-354		M00042797D:D10	CH16COP
865	448446	RTA00002690F 6.09.3.1 .Sec	·	M00040174D:G03	CHISEDT
866	402313	RTA00002686F.f.18.1.P.Sec	·	M0003937+C:H02	CHIZEDT
867	273151	RTA00002685F.c.05.2.P.Sec		M00039951B:B12	CH14EDT
868	404172	RTA00002687F.d.17.2.P.Sec	1	M00043637C:H01	CH20COHLV
869	263630	RTA00002694F.e.10.1.P.Sec	<u> </u>	M00039951B:C03	CH14EDT
870	404277	RTA00002687F.d.18.1.P.Sec	`\	M00039948.A.E03	CH14EDT
871	403557	RTA00002687F.d.10.1.P.Se		M00039319B:H12	CH09LNL
872	375161	RTA00002676F.m.24.2.P.Se	91	M00039135D:G02	CH09LNL
873	376829	RTA00002674F.f.21.1.P.Se	7	M00038639D:F07	CH09LNL
874		RTA00002672F.c.02.2.P.Se	7	M00005351C:G05	CH02COH
875		RTA00002709F.a.24.1.P.Se	q F	M00039760B:B08	CHISEDT
876		RTA00002686F.b.17.1.P.Se	q F	M00027212D:E03	CH04MAL
877		RTA00002713F.b.04.1.P.Se	q F	M00003770A:E05	CHOICOH
878		RTA00002661F.e.11.1.P.Se	q F		CH09LNL
879		RTA00002670F.c.05.2.P.Se	:q	M00033345D:A09	CH09LNL
		RTA00002670F.c.08.2.P.Se	q F	M00033346C:A05	CH12EDT
880		RTA00002685F.n.17.2.P.Sc	eq F	M00039609D:F07	CHOCCOH
38		RTA00002662F.b.23.2.P.S	eq F	M00006712C:H09	CHI-EDT
883		RTA00002688F.d.21.2.P.S	eq F	M00040394A:D04	CH04MAL
88:		RTA00002664F.c.18.2.P.S	eq F	M00026918B:D01	
884		RTA00002689F.b.11.3.P.S	eq F	M00042560A:F12	CHISCON
88		RTA00002682F.d.17.1.P.S		M00039936C:C05	CH09LNL
88		RTA00002664F.d.21.2.P.S	<u> </u>	M00027035D:C06	CH04MAL
38		K1A00002604F.d.21.2.F.S		M00042534A:A05	CH15CON
88		RTA00002689F.3.12.1.P.S		M00032545B:H09	CH08LNH
88		RTA00002666F.c.24.1.P.S		M00027818C:C07	CH04MAL
89		RTA00002714F.b.20.1.P.S		M00040377C:G07	CHITEDL
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901	192535	RTA00002663F.m.14.1.P.Seq	F	M00039820B:B06	CH09LNL
902	377926	RTA00002680F.1.16.2.P.Seq	F	M00039820B:B00	CH04MAL
903	186055	RTA00002712F.i.11.1.P.Seq	F	M00040129D:E10	CH09LNL
904	380498	RTA00002684F.f.11.2.P.Seq	F	M00040129D:E10	CH12EDT
905	400236	RTA00002685F.i.18.2.P.Seq	F		CH14EDT
906	401070	RTA00002688F.d.12.2.P.Seq	. F	M00040390A:H02	CHISCON
907	452622	RTA00002692F.b.14.2.P.Seq	F	M00042962D:C05	CH18CON
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913	403242	RTA00002693F.a.07.2.P.Seq	F	M00042614B:B05	CH19COP
914	453313	RTA00002692F.f.11.2.P.Seq	F	M00043067D:D10	CH18CON
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920	236906	RTA00002687F.e.20.2.P.See		M00039958C:B09	CH14EDT
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925	403578	RTA00002687F.i.01.2.P.Se		M00040296D:E09	CH14EDT
926	403578	RTA00002687F.h.24.2.P.Se	71	M00040294D:D12	CH14EDT
927	403371	RTA00002687F.h.19.2.P.Se		M00043125A:B11	CH18CON
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933		RTA00002687F.f.02.2.P.Se	q F	M00039816C:D05	CH09LNL
934		RTA00002680F.k.19.1.P.S	eq F	M00039810C:D02	CH19COP
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936		RTA00002710F.d.09.1.P.S	eq F		CH20COHL
93		RTA00002694F.b.18.1.P.S	eq F	M00043446C:E12	CH20COHL
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93		RTA00002694F.d.13.1.P.S	eq F	M00043513D:G08	CH20COHL
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944	85388	RTA00002674F.c.06.2.P.Seq	F	M00039124C:H08	CH09LNL
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950	124813	RTA00002685F.j.10.2.P.Seq	F	M00039564B:C01	CH12EDT
951	454627	RTA00002693F.f.09.2.P.Seq	F	M00043210C:E05	CH19COP
952	169464	RTA00002663F.i.19.1.P.Seq	F	M00022602.4:E09	CH03MAH
953	451654	RTA00002692F.f.02.2.P.Seq	F	M00043044D:A09	CH18CON
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956	450845	RTA00002691F.f.10.1.P.Seq	F	M00043410C::A09	CH17COHLV
957	448177	RTA00002690F.e.12.1.P.Seq	F	M00042839B:B11	CH16COP
958	402617	RTA00002686F.b.21.1.P.Seq	F	M00040131B:D11	CHISEDT
959	378014	RTA00002680F.g.17.1.P.Seq		M00039799.A:D10	CH09LNL
960	124813	RTA00002685F.j.10.1.P.Seq	F	M00039564B:C01	CH12EDT
961	29450	RTA00002663F.d.07.1.P.Seq		M00022054A:H03	СНОЗМАН
962	400486	RTA00002685F.e.02.1.P.Seq		M00039496B:D08	CH12EDT
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964	448177	RTA00002690F.e.12.2.P.Seq	F	M00042839B:B11	CH16COP
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991	400921	RTA00002685F.b.18.1.P.Seq	F	M00039371B:H06	
992	373874	RTA00002672F.c.22.2.P.Seq	F	M00038663D:H10	CH09LNL
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994	453237	RTA00002693F.c.02.2.P.Seq	F	M00043108A:F06	
995	449294	RTA00002690F.c.13.3.P.Seq	F	M00042770C:C04	CH16COP
996	404260	RTA00002687F.c.11.1.P.Seq	F	M00039942D:C01	CHI4EDT
997	378014	RTA00002680F.g.17.2.P.Seq	F	M00039799A:D10	CH09LNL
998	404726	RTA00002688F.a.18.2.P.Seq	F	M00040371C:H05	CH14EDT
999	451347	RTA00002691F.b.11.3.P.Seq	F	M00043311C:E03	CH17COHLV
1000	401154	RTA00002685F.e.06.2.P.Seq	F	M00039497C:C06	CH12EDT
1001	401870	RTA00002686F.b.22.1.P.Seq	F	M00040131C:F03	CHISEDT
.1002	400170	RTA00002685F.b.03.2.P.Seq	F	M00039366C:B07	CH12EDT
1003	25387	RTA00002711F.f.19.1.P.Seq	F	M00023001C:C08	CH03MAH
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1006	372930	RTA00002670F.j.12.2.P.Seq	F	M00033437C:A07	CH09LNL
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1008	403397	RTA00002687F.h.02.2.P.Seq	F	M00040219B:D02	CHI4EDT
1009	449337	RTA00002690F.c.18.3.P.Seq	F	M00042774C:C05	CH16COP
1010	403561	RTA00002688F.d.06.2.P.Seq	F	M0004038 C: E07	CH14EDT
1011	134182	RTA00002692F.d.13.2.P.Seq	F	M00043011A:H12	CHISCON
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1013	376138	RTA00002674F.m.05.1.P.Seq		M00039169A:E12	CH09LNL
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1024	401553	RTA00002685F.d.08.2.P.Seq	F	M00039482B:G02	CHIZEDT
1025	451092	RT.A00002691F.d.17.3.P.Seq	F	M00043377A:C03	CH17COHLV
1026	403978	RTA00002687F.g.09.2.P.Seq		M00040208B:A07	CH14EDT
1027	377186	RTA00002682F.m.07.1.P.Seq		M0004001+D:F03	CH09LNL
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1040	406092	RTA00002685F.k.11.1.P.Seq	F	M00039584C:C11	CH12EDT
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	402339	RTA00002686F.o.16.1.P.Seq		M00040282.4:.403	CHISEDT
1048	401766	RTA00002686F.g.14.1.P.Seq		M00040181D:H10	CHISEDT
1049	449669	RTA00002690F.c.10.3.P.Seq	F	M00042767B:G10	CH16COP
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1052	403868	RTA00002687F.1.05.1.P.Seq		M00040323B:C12	CH14EDT
1053	403242	RTA00002686F.f.16.1.P.Seq		M00040174C:E10	CHISEDT
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1067		RTA00002686F.f.05.1.P.Sec	<u>' </u>	M00040169B:F08	CHISEDT
1068		RTA00002685F.o.23.2.P.Sec		M00039642D:H09	CHIZEDT
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1071		RTA00002690F.c.22.3.P.Sec		M00042781A:A07	CH16COP
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1074		RTA00002694F.e.05.1.P.Sec		M00043632D:F09	CH20COHLV
1075		RTA00002685F.o.04.1.P.Sec		M00039625B:G08	CHIZEDT
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1077		RTA00002686F.g.20.1.P.Sec		M00040184C:A11	CHISEDT
1078		RTA00002676F.n.01.2.P.Se		M00039319B:H12	CH09LNL
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1080		RTA00002685F.a.14.2.P.Set	<u> </u>	M00039361B:E01	CHIZEDT
108	400567	KTAUUCULUUJT a. 14.2.1. See	<u> </u>		

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1084	400450	RTA00002685F.j.22.1.P.Seq	F	M00039570.A:D10	CH12EDT
1085	375373	RTA00002676F.h.12.1.P.Seq	F	M00039300C:C09	CH09LNL
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1139	449832	RTA00002691F.e.13.1.P.Seq		M00040093B:C02	CH09LNL
1140	379004	RTA00002683F.n.09.2.P.Seq	F	M00043430B:C02	CH20COHLV
1141	455211	RTA00002694F.b.07.1.P.Seq	F	M00040093D:D03	CH09LNL
1142	379021	RTA00002683F.n.13.2.P.Seq	F	M00039785D:G05	CH09LNL
1143	376279	RTA00002680F.d.10.2.P.Seq	F	M00039783D:G03	CH09LNL
1144	374373	RTA00002681F.n.21.1.P.Seq	F	M00039903A:H07	CHISEDT
1145	97668	RTA00002686F.d.19.1.P.Seq	F		CH12EDT
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1179	18139	RTA00002708F.f.10.1.P.Seq	F	M00039746C:H06	CH14EDT
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1183	400973	RTA00002685F.c.06.2.P.Seq	F	M00043405A:D11	CH17COHLV
1184	450198	RTA00002691F.e.23.2.P.Seq	F	M00043406B:G12	CH17COHLV
1185	451502	RTA00002691F.f.03.2.P.Seq	F	M00043220B:C04	CH19COP
1186	454414	RTA00002693F.f.18.2.P.Seq	F	M00043081D:F05	CH19COP
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1138	403700	RTA00002687F.g.03.1.P.Seq		M00040294D:D12	CH14EDT
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1203	271092	RTA00002690F.b.23.2.P.Sec	F	M00039517B:G12	CHIZEDT
1204	400864	RTA00002685F.g.17.2.P.Sec	q F	M00032876C:D06	CH08LNH
1205	235855	RTA00002667F.o.06.1.P.Sec	q F	M00040183A:F07	CHISEDT
1206	402789	RTA00002686F.g.16.1.P.Se	q F	M00022467C:B12	CH03MAH
1207	19826	RTA00002710F.k.05.1.P.Se	q F	M00039984D:G12	CH09LNL
1208	380157	RTA00002682F.h.19.1.P.Se	q F	M00039500C:C04	CH12EDT
1209	401187	RTA00002685F.e.15.2.P.Se	q F	M00039300C:C04	CH08LNH
1210	427346	RTA00002665F.b.01.3.P.Se	q F	M00040138B:H03	CHISEDT
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1225	403541	RTA00002687F.p.20.1.P.Seq	F	M00043385D:A02	CH17COHLV
1226	450773	RTA00002691F.d.24.3.P.Seq	F	M00039595C:E05	CH12EDT
1227	376236	RTA00002685F.1.24.2.P.Seq	F	M00040385C:D02	CH14EDT
1228	422357	RTA00002688F.c.21.1.P.Seq	F	M00040351B:F02	CH14EDT
1229	404532	RTA00002687F.p.10.2.P.Seq	F	M00040317D:F02	CHI4EDT
1230	403693	RTA00002687F.j.23.1.P.Seq	F	M00040317D:F02	CH14EDT
1231	403693	RTA00002687F.j.23.2.P.Seq		M00039624B:F12	CH12EDT
1232	401515	RTA00002685F.o.02.2.P.Seq	F	M00040351B:F02	CH14EDT
1233	404532	RTA00002687F.p.10.1.P.Seq	F	M00043002A:E05	CH18CON
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1235	18003	RTA00002711F.b.04.1.P.Seq	F	M00039973 D:C08	CH09LNL
1236	377014	RTA00002682F.f.13.1.P.Seq	F	M00039973D:C00	CH14EDT
1237	404232	RTA00002687F.n.12.2.P.Seq	F		CH14EDT
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1239	406263	RTA00002685F.d.14.1.P.Seq	F	M00039493.A:C04	CHISCON
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125		RTA00002689F.b.14.1.P.Se	q F	M00042566C.C05	CH15CON
125		RTA00002677F.k.12.2.P.Se	g F	M00039411C:E07	CH09LNL
125		RTA00002678F.e.10.2.P.Se	a F	M00039458B:H11	CH09LNL
125		RTA00002678F.f.10.2.P.Se	a F	M00043066B.H11	CH18CON
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126		RTA00002670F.n.24.2.P.Se	q F	M00039855C.F01	CH09LNL
126	379602	RTA00002681F.c.21.2.P.Sc		M00039746C:H05	CH14EDT
126	403896	RTA00002687F.a.04.1.P.Sc		M00040219B:D02	CHITEDT
126		RTA00002687F.h.02.1.P.S	eq F	M00039755A:B08	CHISEDI
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1273	428491	RTA00002666F.c.05.1.P.Seq	F	M00032535D:H01	CH08LNH
1274	54656	RTA00002661F.i.22.2.P.Seq	F	M00004372B:F07	CH01COH
1275	379183	RTA00002679F.i.17.1.P.Seq	F	M00039688C.G06	CH09LNL
1276	25594	RTA00002711F.f.07.1.P.Seq	F	M00022968B:E02	CH03MAH
1277	403355	RTA00002687F.d.11.1.P.Seq	F	M00039948D:D11	CH14EDT
1278	16789	RTA00002709F.b.09.2.P.Seq	F	M00005382B:F08	CH02COH
1279	23292	RTA00002708F.c.02.1.P.Seq	F	M00003750D:E06	CH01COH
1280	373982	RTA00002673F.b.24.2.P.Seq	F	M00039058A:A04	CH09LNL
1281	373982	RTA00002673F.c.01.2.P.Seq	F	M00039058A::A04	CH09LNL
1282	449911	RTA00002691F.e.02.2.P.Seq	F	M00043384B:B02	CH17COHLV
1283	450633	RTA00002691F.f.02.2.P.Seq	F	M00043405C:G12	CH17COHLV
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1359	456629	RTA00002694F.d.04.1.P.Sec		M00033218A:C04	CH08LNH
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1366	451438	RTA00002691F.d.23.3.P.Seq	F	M00043383C:F12	CH17COHLV
1367	379011	RTA00002681F.n.23.1.P.Seq	F .	M00039903C:D01	CH09LNL
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1375	450627	RTA00002691F.f.01.2.P.Seq	F	M00043405C:G02	CH17COHLV
1376	375589	RTA00002680F.f.06.2.P.Seq	F	M00039794.4:E04	CH09LNL
1377	379011	RTA00002681F.n.23.2.P.Seq	F	M00039903C:D01	CH09LNL
1378	16789	RTA00002709F.b.09.1.P.Seq	F	M00005382B:F08	CH02COH
1379	427346	RTA00002665F.a.24.3.P.Seq	F	M00028066C:D07	CH08LNH
1380	49540	RTA00002712F.e.01.1.P.Seq	F	M00023399C:E10	CH04MAL
1381	14440	RTA00002674F.e.14.2.P.Seq	F	M00039129C:D04	CH09LNL
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1383	43782	RTA00002662F.d.21.2.P.Seq	F	M00007165B:G11	СН02СОН
1384	212635	RTA00002666F.p.01.1.P.Seq	F	M00032688D:DII	CH08LNH
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1386	18501	RT.A00002669F.g.23.3.P.Seq	F	M00033217B:H07	CH08LNH
1387	400310	RTA00002688F.b.05.2.P.Seq	F	M00040375C:B06	CH14EDT
1388	403796	RTA00002687F.h.17.1.P.Seq	F	M00040293 D:G04	CH14EDT
1389	452314	RTA00002694F.a.21.1.P.Seq	F	M00043416C:A02	CH20COHLV
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1392	450523	RTA00002691F.e.19.2.P.Seq	F	M00043401D:G08	CH17COHLV
1393	289535	RTA00002693F.f.06.1.P.Seq	F	M00043202B:F01	CH19COP
1394	374736	RTA00002673F.o.08.2.P.Seq	F	M00039112B:C05	CH09LNL
1395	378912	RTA00002672F.n.01.2.P.Seq	F	M00039036C:B05	CH09LNL
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1397	372811	RTA00002670F.c.12.2.P.Seq	F	M0003334~C:F02	CH09LNL
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1407	431066	RTA00002669F.c.17.3.P.Seq	F	M00033189D:F08	CH08LNH
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1414	375589	RTA00002680F.f.06.1.P.Seq	F	M00039794A:E04	CH09LNL
1415	375789	RTA00002674F.a.16.1.P.Seq	F	M00039120C:H03	CH09LNL
1416	456227	RTA00002694F.c.16.1.P.Seq	F	M00043465C:C09	CH20COHLV
1417	455852	RTA00002694F.a.02.1.P.Seq	F	M00042592A:H10	CH20COHLV
1418	25169	RTA00002710F.m.05.1.P.Seq	F	M00022579C:C11	CH03MAH
1419	376524	RTA00002678F.h.23.2.P.Seq	F	M00039477A:B03	CH09LNL
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1428	418358	RTA00002686F.m.07.1.P.Seq	F	M00040265D:B07	CHISEDT
1429	380263	RTA00002689F.a.22.1.P.Seq	F	M00042543C:G04	CHISCON
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1431	451679	RTA00002693F.a.04.2.P.Seq	F	M00042612D:F06	CH19COP
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1435	402494	RTA00002686F.h.16.1.P.Seq	F	M00040191A:B09	CHI3EDT
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1462	74344	RTA00002661F.f.10.1.P.Seq	F	M00003902.4:C03	CH01COH
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1469	376366	RT.A00002677F.h.05.2.P.Seq	F	M00039397B:H09	CH09LNL
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1473	379721	RT.A00002676F.b.20.2.P.Seq	F	M00039276B:H09	CH09LNL
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1482	449080	RTA00002690F.a.04.2.P.Seq	F	M0004234TD:H11	CH16COP
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1676	380596	RTA00002671F.a.03.1.P.Seq	F	M00033584D:G11	CH09LNL
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1693	380354	RTA00002670F.n.23.2.P.Seq	F	M00033570B:C08	CH02COII
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1747	38891	RTA00002677F.h.24.2.P.Seq	F	M00039401B:D02	CH0ºLNL
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1806	121871	RTA00002713F.a.09.1.P.Seq	F	M00027198B:B08	
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1815	119478	RTA00002686F.n.07.1.P.Seq	F	M00040271C:D08	CHISEDT
1816	403189	RTA00002687F.g.16.2.P.Seq	F	M00040217D:B07	CH!+EDT
1817	129692	RTA00002679F.e.13.1.P.Seq	F	M00039673A:F09	CH09LNL
1818	86668	RTA00002664F.a.10.2.P.Seq	F	M00023352B:F03	CH04MAL
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1836	429663	RTA00002667F.m.21.1.P.Seq	F	M00032864B:B09	CH08LNH
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4	10924	RTA00002903F.I.10.1.P.Seq	F	M00007037D:D10	CH02COH
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28	1870	RT.400002910F.m.0S.1.P.Seg	<u> </u>	M00023020C:H03	CH03MAH
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554	186084	RTA00002912F.f.18.1.P.Seq	F	M00039805B:B06	CH09LNL
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625	186324	RTA00002912F.d.17.1.P.Seq		M00027274A:A09	CH01COH
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637	17237	RTA00002901F.1.12.1.P.Seq	F	M100005616B:F07 M00005346D:A03	CH02COH
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650	7870	RTA00002905F.b.22.1.P.Sec	ıl F	M00007973B:D11	CHOUNTAL

SEQ		CEO NAME	OD TEATE ATTOM	CLONEID	I TOO VOV
ID	CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
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704	186729	RTA00002911F.d.19.2.P.Seq	F	M00026850B:C09	CH04MAL
705	33658	RT.A00002886F.j.07.1.P.Seq	F	M00001361B:A12	CH01COH
706	186755	RTA00002912F.i.18.1.P.Seq	F	M00027400D:H02	CH04MAL
707	4262	RTA00002897F.a.04.1.P.Seq	F	M00004208.4:D08	CH01COH
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717	25320	RTA00002910F.e.19.1.P.Seq	F	M00022857B:A09	CH03MAH
718	4924	RTA00002930F.g.01.2.P.Seq	F	M00055805.A:H02	CH15CON
719	21170	RTA00002900F.1.13.1.P.Seq	F	M00005365.A:F05	CH02COH
720	9258	RT.A00002890F.h.17.1.P.Seq	F	M00001618C:D01	CH01COH
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733	12595	RTA00002904F.c.06.1.P.Seq		M00007197B:B05	CH02COH
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757	6078	RTA00002930F.c.11.1.P.Seq	F	M00039377B:E05	CH12EDT
758	12543	RTA00002927F.b.14.1.P.Seq	F	M00055794A:E10	CH15CON
759	9686	RTA00002930F.f.19.1.P.Seq	F	M00042732B:H06	CH15CON
760	3369	RTA00002930F.b.12.1.P.Seq	F	M00004087C:E02	CHOICOH
761	6891	RTA00002895F.i.03.1.P.Seq	F	M00003822C:A09	CHOICOH
762	13666	RTA00002892F.i.05.1.P.Seq	F	M000056458C:E01	CHISCON
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766	1596	RTA00002922F.m.18.1.P.Seq	F	M00039113D:1112 M00039411D:D09	CH09LNL
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SEQ	a	SEQ NAME	ORIENTATION	CLONE ID	LIBR.ARY
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827	3441	RTA00002935F.i.13.1.P.Seq			CH15CON_
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926	9484	RTA00002934F.a.13.1.P.Sec		M00001633D:C11	CHOICOH
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17636 2328 15414 11948 24759 15152 14917 12941 29676 17789 45097 18407 22309 109382 92273	RTA00002933F.c.19.1.P.Seq RTA00002935F.e.05.1.P.Seq RTA00002935F.p.13.1.P.Seq RTA00002895F.o.01.1.P.Seq RTA00002903F.n.05.1.P.Seq RTA00002925F.g.01.1.P.Seq RTA00002922F.b.02.1.P.Seq RTA00002889F.c.15.1.P.Seq RTA00002931F.b.03.1.P.Seq RTA00002931F.b.03.1.P.Seq RTA00002909F.b.11.1.P.Seq RTA00002909F.b.11.1.P.Seq	F F F F F F F F F	M00043222C:B06 M00054579A:C02 M00055423C:H10 M00004118C:D12 M00007082D:E05 M00039873B:H04 M00038616D:B07 M00001532A:G08 M00042788A:F04 M00001671A:H10 M00040247D:D02 M100022546B:E05	CH19COP CH17COHLV CH17COHLV CH01COH CH02COH CH09LNL CH09LNL CH01COH CH16COP CH01COH CH13EDT CH03MAH
2328 15414 11948 24759 15152 14917 12941 29676 17789 45097 18407 22309 109382 92273	RTA00002935F.e.05.1.P.Seq RTA00002935F.p.13.1.P.Seq RTA00002895F.o.01.1.P.Seq RTA00002903F.n.05.1.P.Seq RTA00002925F.g.01.1.P.Seq RTA00002922F.b.02.1.P.Seq RTA00002889F.c.15.1.P.Seq RTA00002931F.b.03.1.P.Seq RTA00002931F.b.03.1.P.Seq RTA00002909F.b.11.1.P.Seq RTA00002909F.b.11.1.P.Seq	F F F F F F F F	M00054579 A:C02 M00055423C:H10 M00004118C:D12 M00007082D:E05 M00039873B:H04 M00038616D:B07 M00001532 A:G08 M00042788 A:F04 M00001671 A:H10 M00040247 D:D02 M00022546B:E05	CH17COHLV CH17COHLV CH01COH CH02COH CH09LNL CH09LNL CH01COH CH16COP CH01COH CH13EDT CH03MAH
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24759 15152 14917 12941 29676 17789 45097 18407 22309 109382 92273	RTA00002903F.n.05.1.P.Seq RTA00002925F.g.01.1.P.Seq RTA00002922F.b.02.1.P.Seq RTA00002889F.c.15.1.P.Seq RTA00002931F.b.03.1.P.Seq RTA00002891F.a.21.1.P.Seq RTA00002928F.g.06.1.P.Seq RTA00002909F.b.11.1.P.Seq RTA00002900F.n.19.1.P.Seq	F F F F F F F	M00007082D:E05 M00039873B:H04 M00038616D:B07 M00001532A:G08 M00042788A:F04 M00001671A:H10 M00040247D:D02 M100022546B:E05	CH02COH CH09LNL CH09LNL CH01COH CH16COP CH01COH CH13EDT CH03MAH
15152 14917 12941 29676 17789 45097 18407 22309 109382 92273	RTA00002925F.g.01.1.P.Seq RTA00002922F.b.02.1.P.Seq RTA00002889F.c.15.1.P.Seq RTA00002931F.b.03.1.P.Seq RTA00002891F.a.21.1.P.Seq RTA00002928F.g.06.1.P.Seq RTA00002909F.b.11.1.P.Seq RTA00002900F.n.19.1.P.Seq RTA00002907F.k.13.1.P.Seq	F F F F F F	M00039873B:H04 M00038616D:B07 M00001532A:G08 M00042788A:F04 M00001671A:H10 M00040247D:D02 M100022546B:E05	CH09LNL CH09LNL CH01COH CH16COP CH01COH CH13EDT CH03MAH
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12941 29676 17789 45097 18407 22309 109382 92273	RTA00002889F.c.15.1.P.Seq RTA00002931F.b.03.1.P.Seq RTA00002891F.a.21.1.P.Seq RTA00002928F.g.06.1.P.Seq RTA00002909F.b.11.1.P.Seq RTA00002900F.n.19.1.P.Seq RTA00002907F.k.13.1.P.Seq	F F F F F	M00001532 A: G08 M00042788 A: F04 M00001671 A: H10 M00040247 D: D02 M100022546B: E05	CH16COP CH01COH CH13EDT CH03MAH
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SEQ		250 24 255	ORIENTATION	CLONE ID	LIBRARY
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1253 1254 1255 1256 1257 1258 1259 1260	12315 4930 12018 10501 11314 6426 2205 6991	RT.A00002907F.1.24.2.P.Seq RT.A00002930F.g.04.2.P.Seq RT.A00002924F.c.16.1.P.Seq RT.A00002930F.h.23.1.P.Seq RT.A00002935F.k.03.1.P.Seq	F F F	M00022240D:B11 M00055810C:D03	CH03MAH CH15CON
1254 1255 1256 1257 1258 1259 1260	4930 12018 10501 11314 6426 2205 6991	RTA00002930F.g.04.2.P.Seq RTA00002924F.c.16.1.P.Seq RTA00002930F.h.23.1.P.Seq RTA00002935F.k.03.1.P.Seq	F F F	M00055810C:D03	CH15CON
1255 1256 1257 1258 1259 1260	12018 10501 11314 6426 2205 6991	RTA00002924F.c.16.1.P.Seq RTA00002930F.h.23.1.P.Seq RTA00002935F.k.03.1.P.Seq	F F		
1256 1257 1258 1259 1260	10501 11314 6426 2205 6991	RTA00002930F.h.23.1.P.Seq RTA00002935F.k.03.1.P.Seq	F	M00039438B:D08	י די יסטוזט י
1256 1257 1258 1259 1260	10501 11314 6426 2205 6991	RTA00002935F.k.03.1.P.Seq			CH09LNL
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1316	186798	RTA00002911F.f.11.1.P.Seq		M00001369A:G06	CH01COH
1317	13193	RTA00002886F.1.16.1.P.Seq		M00033149B:E10	CH08LNH
1318	3554	RTA00002919F.i.14.1.P.Seq		M00022446C:H06	CH03MAH
1319	19991	RT.A00002908F.h.11.1.P.Seq		M00005703D:G10	CH02COH
1320	173046	RTA00002901F.o.19.1.P.Seq		M00043016B:F09	CH18CON
1321	21798	RTA00002932F.b.12.1.P.Seq		M00004359A:E01	CH01COH_
1322	11303	RTA00002898F.i.11.1.P.Seq		M00032494C:H08	CH08LNH
1323	4026	RTA00002915F.m.02.2.P.Sec	F	M0003245-0:1105	СНОЗМАН
1324	94859	RTA00002909F.i.23.1.P.Seq	·F	M00022240D:B11	CH03MAH
1325	12315	RTA00002907F.m.01.1.P.Sec	F	M00022240D:B11	CH03MAH
1326	4822	RTA00002909F.1.16.1.P.Sec	F	M00022684A:E06	СНОЗМАН
1327	97129	RTA00002909F.1.13.1.P.Sec	F	M00022004A:E00	CH01COH
1328	15996	RTA00002897F.1.09.1.P.Sec	F	M00004231A:C04	CH08LNH
1329		RTA00002918F.c.01.1.P.Sec	F	M00032833B3C04	CH02COH
1330		RTA00002902F.h.08.1.P.Sec	F	MI00006646A:A07	CH02COH
1331	15642	RTA00002902F.g.06.1.P.Sec	9 ·F 、	M00000046A:A07	CH08LNH
1332	· · · · · · · · · · · · · · · · · · ·	RTA00002916F.f.05.1.P.Sec	F	M00032567B:G05	CH02COH
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1335		.RTA00002916F.b.19.1.P.Se	q <u>F</u>	M00032541C:G03	CH04MAL
1336		RTA00002911F.e.24.1.P.Se	q F	M00026900 A:H07	CH03MAH
1337		RTA00002907F.o.19.1.P.Se	q F	M100022273A:E03	CH01COH
1338		RTA00002887F.a.09.1.P.Se	q F	M00001385A:E07	CH08LNH
		RTA00002914F.h.23.1.P.Se	q F	M00028212D:C05	
1339		RTA00002917F.g.15.1.P.Se	q F	M00032727A:E04	CH08LNH
1340		RT.A00002923F.o.07.1.P.Se	g F	M00039381C:C07	CH09LNL
1341		RT.A00002928F.d.02.1.P.Se	q F	M100040169.A:G06	CHISEDT
1347		RTA00002892F.f.10.2.P.Se		M00003814A:G05	CH01COH
1343		RTA00002919F.f.14.1.P.Se	g F	M00033072A:A09	CH08LNH
134		RTA00002888F.a.04.1.P.Se		M00001433B:E02	CH01COH
134		RTA00002897F.d.03.1.P.Sc		M00004225D:E03	CH01COH
134		RTA00002897F.d.051FF.Sc RTA00002935F.k.11.1.P.Sc		M00055055C:F01	CH17COHL
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134		RTA00002911F.1.01.117.50		M00033434D:F05	CH09LNL
134		RTA00002921F.g.24.1.F.S		M000224~4B:C08	
135	0 25844	K1A00002903F.k.23.1.F.3	-41		

				 	
SEQ			OPTIVITATION	CLONE ID	LIBRARY
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1352	7695	RT.A00002892F.n.18.2.P.Seq	F	M00003845A:C07	CHOICOH
1353	16997	RTA00002922F.k.15.1.P.Seq	F	M0003910 A: E12	CH09LNL CH03MAH
1354	25441	RTA00002906F.i.08.1.P.Seq	F	M00021981A:C02	
1355	4303	RTA00002897F.o.20.1.P.Seq	· F	M00004295D:C07	CHOICOH
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1358	11766	RTA00002925F.f.20.1.P.Seq	F	M00039871C:G05	CH09LNL CH01COH
1359	13618	RTA00002893F.o.15.1.P.Seq	F	M00003963D:F01	CH09LNL
1360	13903	RTA00002923F.c.18.1.P.Seq	F	M00039204.A:E09	
1361	10673	RTA00002927F.h.23.1.P.Seq	F	M100039646A:E06	CH12EDT
1362	17412	RTA00002932F.b.11.1.P.Seq	F	M00043015D:D05	CH18CON
1363	2218	RTA00002919F.a.20.1.P.Seq	F	M00033028C::A02	CH08LNH
1364	5858	RTA00002923F.i.01.1.P.Seq	F	M00039275B:E02	CH09LNL
1365	2510	RTA00002898F.b.14.1.P.Seq	F	M00004316.A:B03	CH01COH
1366	8050	RTA00002900F.n.04.1.P.Seq	F	M00005383.A:C11	CH02COH
1367	186538	RTA00002929F.e.18.1.P.Seq	F	M00040329.A:H05	CH14EDT
1368	25427	RT.A00002935F.n.20.1.P.Seq	F	M00055337B:C04	CH17COHLV
1369	24098	RT.A00002901F.a.10.1.P.Seq	F	M00005422D:H02	CH02COH
1370	123823	RTA00002905F.h.08.1.P.Seq	F	M00008071D:H03	CH03MAH
1371	3644	RTA00002901F.c.03.1.P.Seq	F	M00005445D:D04	CH02COH
1372	27783	RTA00002917F.a.17.1.P.Seq	F	M00032666.A:C02	CH08LNH
1373	1682	RTA00002910F.b.03.1.P.Seq	F	M100022801D:D09	CH03M.AH
1374	3200	RT.A00002887F.e.07.1.P.Seq		M00001393C:F04	CH01COH
1375	8442	RTA00002917F.h.23.1.P.Seq		M0003273-B:E12	CH08LNH
1376	15353	RTA00002910F.e.11.1.P.Seq		M0002285÷C:G07	CH03MAH
1377	6314	RTA00002922F.b.06.1.P.Seq		M0003861SD:D08	CH09LNL
1378	93549	RTA00002909F.j.14.1.P.Seq	F	M00022662C:H04	CH03MAH
1379	15496	RTA00002906F.p.03.1.P.Seq	F	M000220SSB:H02	CH03MAH
1380	16572	RTA00002886F.k.03.1.P.Seq		M0000136-A:C09	CH01COH
1381	74821	RTA00002890F.p.21.1.P.Seq		M00001663A:A12	CH01COH
1382	11315	RTA00002S89F.d.12.1.P.Seq		M00001535B:B10	CH01COH
		RTA00002S94F.c.18.1.P.Seq		M00003980D:C06	CH01COH
1383	10859	RTA00002914F.f.04.1.P.Seq		M00028193B:E07	CH08LNH
1384	15391	RTA00002896F.b.18.1.P.Seq		M00004141B:F08	CH01COH
1385	23172	RTA00002S36F.1.05.1.P.Seq		M00001368.A:C02	CH01COH
1386	22510	RTA00002934F.a.08.1.P.Seq		M00043455B:C08	CH20COHLV
1387	17156	RTA00002396F.o.18.1.P.Seq		M00004200C:A04	CHOICOH
1388		RTA00002901F.m.08.1.P.Seq		M00005626D:G11	CH02COH
1389		RTA00002933F.c.11.1.P.Seq		M00043213.A:D05	CH19COP
1390				M0002210:C:D05	СН03М.АН
1391		RTA00002907F.a.18.1.P.Seq		M000333+:-A:B06	CH09LNL
1392		RTA00002921F.c.07.1.P.Seq		M00004365C:G11	CHOICOH
1393		RTA00002S98F.j.11.1.P.Seq		M00004365 A:F11	CHOICOH
1394		RTA00002S97F.i.20.1.P.Seq		M00004269.A:F11	CH08LNH_
1395		RTA00002916F.d.12.1.P.Seq		M0000501-B:F02	CH02COH
1396		RTA00002900F.h.07.1.P.Seq			CH03MAH
1397		RTA00002905F.c.13.1.P.Seq		M00007981C:F07	CH03MAH_
1398	29703	RTA00002907F.d.24.1.P.Seq		M0002214-C:E12	CH04MAL
1399	6811	RTA00002913F.b.07.1.P.Seq		M0002772-D:D04	CH03MAH
1400	12657	RTA00002906F.b.20.1.P.Seq	F	M00021866C:H08	CHOSNIATI

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SEQ		SEQ NAME	ORIENTATION	CLONE ID	LIBR.ARY
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1402	24229	RTA00002920F.b.04.1.P.Seq	F,	M00001338C:F05	CHOICOH
1403	2066-1	RTA00002886F.a.07.1.P.Seq	F	M00006641B:F05	CH02COH
1404	3656	RTA00002902F.f.20.1.P.Seq	F	M00042878D:G06	CH16COP
1405	10998	RTA00002931F.c.07.1.P.Seq	F	M00039081B:G07	CH09LNL
1406	1150	RTA00002922F.j.14.1.P.Seq	F	M00005013D:H05	CH02COH
1407	45221	RTA00002900F.h.06.1.P.Seq	F	M00005423C:A10	CH02COH
1408	34505	RT.A00002901F.a.16.1.P.Seq	F	M00039472B:E05	CH09LNL
1409	8175	RTA00002924F.f.01.1.P.Seq	F	M00039472B:E05	CH09LNL
1410	8175	RTA00002924F.e.24.1.P.Seq	F	M00007081B:C08	CH02COH
1411	19375	RTA00002903F.n.02.1.P.Seq	F	M00040219B:B07	CH14EDT
1412	10866	RTA00002929F.c.15.1.P.Seq	F	M00003763A:B02	CH01COH
1413	24166	RTA00002891F.k.07.1.P.Seq	F	M000037637112C:G12	CH01COH
1414	15333	RTA00002888F.c.12.1.P.Seq		M00001442C:012	СН03МАН
1415	44436	RTA00002907F.b.17.1.P.Seq		M00042560C:G06	CH15CON
1416	9247	RTA00002930F.a.16.1.P.Seq		M00022430C:C06	СНОЗМАН
1417	12317	RTA00002908F.g.13.1.P.Seq		M00001625D:B04	CH01COH
1418	11968	RT.400002890F.i.24.1.P.Seq		M00022499D:D08	СНОЗМАН
1419	14181	RTA00002908F.n.09.2.P.Seq	F	M00022477C:C01	CH03MAH
1420	15359	RTA00002909F.1.02.1.P.Seq		M0003258-A:D06	CH08LNH
1421	46675	RTA00002916F.h.03.1.P.Seq		M00007019B:E01	CH02COH
1422	24898	RTA00002903F.k.17.1.P.Seq		M00021653A:B02	CH03MAH
1423	156424	RTA00002905F.m.22.1.P.Sec		M00005445A:E07	CH02COH
1424	11996	RTA00002901F.b.24.1.P.Seg		M00005445.A:E07	CH02COH
1425	11996	RT.A00002901F.c.01.1.P.Seq		M00003988D:B01	CH01COH
1426	4784	RTA00002894F.e.20.1.P.Seq		M00028210B:H03	CH08LNH
1427	9120	RTA00002914F.h.10.1.P.Seq		M00001632C:A10	CHOICOH
1428	11295	RTA00002890F.j.15.1.P.Seq		M00004162D:F02	CHOICOH
1429		RT.A00002896F.h.05.1.P.Seq		M0002236 D:G11	CH03MAH
1430	20358	RTA00002908F.b.06.1.P.Sed RTA00002921F.h.01.1.P.Sed		M00033434D:F05	CH09LNL
1431	12823			M00021952B:G06	СН03МАН
1432	147419	RTA00002906F.g.05.1.P.Sec		M00033071D:E08	CH08LNH
1433	12174	RTA00002919F.f.13.1.P.Seq		M00004296B:D03	CH01COH
1434		RT.A00002897F.o.24.1.P.Sec		M00003994A:B10	CH01COH
1435	2325	RTA00002894F.g.07.1.P.Sec RTA00002908F.1.05.1.P.Sec		M00022475D:C07	СНОЗМАН
1436				M00033324B:F04	CH08LNH
1437		RTA00002920F.a.09.1.P.Sec		M00022901.A:C05	CH03MAH
1438		RTA00002910F.g.06.1.P.Sec	1	M00022530B:C04	СН03МАН
1439		RT.A00002909F.a.07.1.P.Sec		M00022992B:G12	СН03М.АН
1440		RTA00002910F.k.22.1.P.Sec		M00022516B:C05	CH03MAH
1441		RT.A00002908F.p.07.1.P.Sec RT.A00002887F.o.06.1.P.Sec		M00001426C:F06	CH01COH
1442		R1.400002887F.0.00.1.F.5ec	F	M00004836B:C02	CH02COH
1443		RT.A00002900F.b.07.1.P.Sec		M00022262A:F06	CH03MAH
1444		RT.A00002907F.n.19.1.P.Sec		M00040079B:F06	CH09LNL
1445		RTA00002926F.c.10.2.P.Sec		M00040173D:A04	CHI3EDT
1446		RT.A00002928F.d.07.1.P.Sec		M00026842B:A01	CH04MAL
1447		RTA00002911F.d.08.2.P.Sec	1	M00001406B:H09	CH01COH
1448		RTA00002887F.j.06.1.P.Sec		M0004307 B:F11	CH19COP
1449		RTA00002933F.a.13.1.P.Sec		M00001551D:H09	CHOICOH
1450	773	RT.A00002889F.j.02.1.P.Sec	il	1710000130131105	<u> </u>

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SEQ			OD WINT A TION	CLONE ID	LIBRARY
ID	CLUSTER	SEQ NAME	ORIENTATION		CH12EDT
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1453	24011	RTA00002924F.c.17.1.P.Seq	F	M00039440C:G06 M00026910B:G06	CH04MAL
1454	5930	RTA00002911F.f.08.1.P.Seq	F	M00026910B:G06 M00005822C:A04	CH02COH
1455	21581	RTA00002902F.c.05.1.P.Seq	F		CH09LNL
1456	3662	RTA00002925F.c.07.1.P.Seq	F	M00039826D:E04 M00042719.A:G08	CH15CON
1457	4873	RTA00002930F.b.05.1.P.Seq	F	M00042719.4.G08 M00004161.4:E08	CHOICOH
1458	11214	RTA00002896F.h.01.1.P.Seq		M00003837C:D10	CHOICOH
1459	22888	RTA00002892F.1.09.1.P.Seq	F	M00003937C:D10	CH09LNL
1460	15490	RTA00002925F.k.08.1.P.Seq		M00039932B:A07 M00021676C:G03	CH03MAH
1461	112819	RTA00002905F.o.13.1.P.Seq	F	M00021676C:G03 M00004179D:A12	CHOICOH
1462	19688	RTA00002896F.1.02.1.P.Seq		M00004179B:R12 M00039138B:G05	CH09LNL
1463	15132	RTA00002922F.n.20.1.P.Seq	F	M00039138B:G03 M00028219B:H05	CH08LNH
1464	25022	RTA00002914F.i.21.1.P.Seq		M00028219B:1103 M00001438A:E01	CHOICOH
1465	16303	RTA00002888F.b.12.1.P.Seq		M000014384:E01	CH01COH
1466	16828	RTA00002897F.b.04.1.P.Seq		M00004214:4:203 M00033296C:C11	CH09LNL
1467	14295	RTA00002921F.a.18.1.P.Seq		M00055725D:D09	CH15CON
1468	1979	RTA00002930F.f.06.1.P.Seq		M00033722D:D09	CHOICOH
1469	36248	RTA00002888F.g.05.1.P.Seq		M00001400C:210	CH09LNL
1470	5676	RTA00002926F.b.22.2.P.Seq		M00001428B:C10	CHOICOH
1471	1239	RT.A00002887F.o.21.1.P.Seq		M00001428B:C10	CH08LNH
1472	7937	RTA00002917F.g.22.1.P.Seq	F	M00032723D.1 01 M00026856B:G03	CH04MAL
1473	4483	RTA00002911F.d.22.2.P.Seq		M00028836B:G05 M00039826B:F09	CH09LNL
1474	7796	RT.A00002925F.c.05.1.P.Seq		M00039820B:109	CH08LNH
1475	17330	RTA00002915F.a.03.1.P.Seg	<u> </u>	M00006631C:A04	CH02COH
1476	25620	RTA00002902F.f.09.1.P.Seq		M00039326.A:G07	CH09LNL
1477	20601	RTA00002923F.1.20.1.P.Seq		M00039258C:C01	CH09LNL
1478	6205	RTA00002923F.g.21.1.P.Sec		M00037236C:C01	CH04MAL
1479	726	RTA00002913F.b.16.1.P.Sec		M00027734B:G12	СН03МАН
1480	104999	RTA00002908F.g.17.1.P.Sec		M00033264B:E06	CH08LNH
1481	30321	RTA00002919F.o.17.1.P.Sec		M000332048:200	CH04MAL
1482	5878	RTA00002913F.a.16.1.P.Sec	<u> </u>	M00021649B:A02	СНОЗМАН
1483	5944	RTA00002905F.m.07.1.P.Sec	<u> </u>	M00021647 A:G05	СНОЗМАН
1484		RTA00002908F.i.21.1.P.Seq	F	M0002245 A:H03	CH17COHLV
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1486		RTA00002918F.a.22.1.P.Sec		M00055254A:H03	CH17COHLV
1487		RTA00002935F.n.01.1.P.Sec		M0003251 C:E10	CHOSLNH
1488		RTA00002915F.o.19.2.P.Sec		M0000378C.A:G01	CHOICOH
1489		RTA00002891F.o.03.1.P.Sec	F F	M0000578C:4:G01	CH02COH
1490	15430	RTA00002900F.g.10.1.P.Sec	1	M00039820B:F06	CH09LNL
1491	5637	RT.A00002925F.b.18.1.P.Sec		M00004246B:H07	CH01COH
1492		RTA00002897F.g.15.1.P.Sec	F F	M00004344A:G11	CH01COH
1493	21826	RT.A00002898F.g.06.1.P.Sec		M00004344 A:G11	CH08LNH
1494	22193	RTA00002919F.i.09.1.P.Sec	F	M00004320C:E07	CH01COH
1495	10720	RT.A00002898F.c.14.1.P.Se		M100040003.A:G10	CH09LNL
1496	22491	RT.A00002925F.m.06.1.P.Se		M0003250°D:G08	CH08LNH
149		RT.A00002915F.n.13.2.P.Se		M0003258cC:B04	CHOSLNH
1498	4953	RTA00002916F.h.11.1.P.Se		M00032388C.B04	CH04M.AL
1499	185567	RT.A00002911F.p.08.1.P.Se		M000271783:A11	CH09LNL
1500	25605	RT.A00002924F.m.22.1.P.Sc	eq F	1 MIOO03971C 3.A01	1

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B	CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
1501		RTA00002906F.m.24.1.P.Seq	F	M00022070B:B04	CH03MAH
1502		RTA00002908F.g.02.1.P.Seq		M00022421A:F12	CH03MAH
1503		RTA00002906F.n.01.1.P.Seq		M00022070B:B04	CH03MAH
1504		RTA00002887F.m.22.1.P.Seq		M00001421B:E07	CH01COH

Table 3

	Nearest N	eighbor (BlastN vs. C	enbank)	Nearest Neighb	or (BlastX vs. Non-Redundant Prot	eins)
EQ	I vedicativ	Cignoor (2)			·	
ID ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION		P VALUE
	<none></none>	<none></none>	∠NONE>	<none></none>	V.101.102	<none></none>
2	<none></none>	<none></none>	<none></none>	<none></none>		<none></none>
- 3	<none></none>	<none></none>	<none></none>	<none></none>	V.10.102	<none></none>
4	<none></none>	<none></none>	<none></none>	<none></none>	2.0	<none></none>
5	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
6	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
7	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
8	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
}	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
		<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
10	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
11	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
12	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
13	<none></none>		<none></none>	<none></none>	<none></none>	<none></none>
14	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
15	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
16	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
17	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
18_	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
19	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
20	<none></none>	<none></none>		<none></none>	<none></none>	<no:ne></no:ne>
21	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
22	<none></none>	<none></none>	<:NONE>	CNOINES	GENOME POLYPROTEIN	
		·			[CONTAINS: RNA REPLICASE; HELICASE: COAT PROTEIN] 2.7.7.48) - apple stem grooving virus	
23	<none></none>	<none></none>	<none></none>	548562	(strain P-209)	9.2
_2,	CAUTAL				EXCISION REPAIR PROTEIN ERCC-6 DNA repair helicase ERCC6 - human >gi 182 IS1 (L04791) excision repair protein	
24	<none></none>	<none></none>	<none></none>	416959	[Homo sapiens] (AB014541) KIAA0641 protein	
					[Homo sapiens]	8 -
25	<none></none>	<none></none>	<none></none>	3327096	(U28741) F35D2.1 gene	
		T	1		product [Caenorhabditis	
				44,000	1.	7.9
26	<none></none>	<none></none>	<none></none>	861293	elegans] (AL031032) extensin-like	
					,	5.5
27	<none></none>	<none></none>	<none></none>	3297821	protein	
					transforming growth factor-beta type III receptor - chicken >gi 511843 (L01121) transforming growth factor-beta	
30	ANONET:	<none></none>	<none></none>	2119692	type III receptor [Gallus gallus]	5.1
28 29		<none></none>	<none></none>	2136028	protein kinuse PRK1 - human	5.0
29	<:NOINE>	ZHOINE -				

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oteins)	BlastX vs. Non-Redundant Prot	Negati Neighbo		·			
1		Nearest Neighb	ank)	ibor (BlastN vs. Ger	. Neigh	Noncort N	
P VALUI	DESCRIPTION	ACCESSION	VALUE	DESCRIPTION	- 1		1
1	F040659) No definition line)E3CRB	י ואכ	CCESSION	A
4.6	F040659) No definition inte						\top
+	und [Caenorhabditis elegans]	2746912	<none></none>	1			十
4.5	F010404) ALR [Homo piens]	2358287		<none></none>	≥	<none></none>	0
.	96048) predicted using	2336267	<none></none>	<none></none>	.	<none></none>	. 1
ì	anafinder: cDNA EST		1			SNOWES	1
is	MRI D65516 comes from this		i i		\.		1
į.	ene; cDNA EST ykl91a5.5				- 1		
1	omes from this gene]		- 1		- 1
4.4	Caenorhabditis elegans]		1		1		- 1
	Y14953) SRCR domain,	3877816	<none></none>	**************************************	- 1		- 1
4.1	Y14953) SRCR dollarin			<none></none>	<u>E> </u>	<none></none>	32 l
	nembrane form 2	4140268	<none></none>	•			
4.0	U51183) transposase [Hydra		anoinz	< NONE>	E>	<none></none>	33
	vulgaris)	1708663					
1 20	(1145958) pistil extensin-like	170000	<none></none>	<none></none>	F.	<none></none>	. I
3.9	protein (Nicotiana alata)				-	CHOILE	34
	GLUCOCORTICOID	1184100	<none></none>	<none></none>	m.	N/ONTS:	_
3.9	RECEPTOR (GR)			4.0.	(5)	<none></none>	35
		121073	<none></none>	<none></none>	_		
an	(U75698) ORF 45; contains an			ANUNC	1 <u>E></u>	<none></none>	36
v	extended acidic domain; EBV		1		- 1		
1	extended delate domain				ţ	ţ	
rusl	BKRF4 homolog [Kaposi's		1		1	1	
ius;	sarcoma-associated herpesviru		1		1]	
2.0	homolog, conserved in other		1		1	1	
	gamma-herpesviruses	1718298				l	
į.	(AF006564) alcohol	1/18298	<none></none>	<none></none>			
1	dehydrogenase [Drosophila			410.12	NES	<none< td=""><td>37</td></none<>	37
1.	persimilis) persimilis)		1			1	
	(AF066071) SP85; PsB	2352538	<none></none>	NONES		1	
1 1.	(AF0000/1) St 65, 152		+	<none></none>	NE>_	<none< td=""><td>38</td></none<>	38
	[Dictyostelium discoideum]	3192897	<none></none>				
1			TONE	<none></none>	NE>	<nont< td=""><td>39</td></nont<>	39
, _{ie}			1			1	ئـــــــــــــــــــــــــــــــــــــ
13	(L33421) This CDS feature i		1	1		Ţ	ļ
auon	included to show the translat	l · .	1	1		1	1
gion.	Let the corresponding V_region	1	1	}		1	l
iers	Presently translation qualifie	l	}	1		1	1
legal	features are lile		ļ	1		ì	
		561645	<none></none>	NONE>_		1	}
}	Conefinder: cDNA EST	Τ -	+	ENOINES	ONE>	O <non< td=""><td>4</td></non<>	4
m this	EMBL:D35016 comes from	1	.1	1			
	DNA EST	1	1	1		1	1
on this	gene; cDNA EST	1	- {			1	1
	EMBL:D32583 comes from	l	1				1
	gene; cDNA EST	1	1			l l	1
om this	EMBL:D35258 comes from	1	1	1		1	١
1	Inner CONA EST	1	l l	}		l	
om this	EMBL:C11471 comes from	1	1	-		1	١
:C	gene: cDNA EST EMBL:C	1	1	1		1	1
	gene; CDNA EST EMBE.	387885	<none< td=""><td><none></none></td><td></td><td>Ţ</td><td>-1</td></none<>	<none></none>		Ţ	-1
-			1 2,70,72	I NONE>	NONE		

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113)	or (BlastX vs. Non-Redundant Protei	Nearest Neighb	abank)			_
			HOMIK)	ghbor (BlastN vs. Ge	Nearest Ne	
VALUE	DESCRIPTION P	COTCCION				Q
	(U75903) UGTIA7 [Rattus	ACCESSION	PVALUE	DESCRIPTION /	CCESSION	
1.0	norvegicus]				+	-
	(AF005370) putative immediate	1658571	<none></none>	<none></none>	<none></none>	2
	early protein [Alcelaphine				-	-
0.86	hamacuirus 11	222024		•	}	- 1
	(AB011167) KIAA0595 protein	2338034	<none></none>	<none></none>	<none></none>	13
0.42	[Homo sapiens] HYPOTHETICAL 92.7 KD	3043714				"
	HYPOTHETICAL 92.7 KD	3043714	<none></none>	<none></none>	<none></none>	44
	PROTEIN IN ASN2-PHB1		1 1			-
	INTERGENIC REGION		1		ļ	ļ
	>gi 2131678 pir S64439		1 1		!	1
	hypothetical protein YGR130c -		1		j	1
•	yeast (Saccharomyces	•	1	•	ì	ı
	cerevisiae)		1 1	•		
	>gi 1323215 gnl PID e243523		1			1
0.40.	(7.72915) ORF YGR130c		1			l
0.40	[Saccharomyces cerevisiae]	1723710	<none></none>	. 0_	1	
	HYPOTHETICAL 92.7 KD		NONE	<none></none>	<none></none>	45
	PROTEIN IN ASN2-PHB1		1			
	INTERGENIC REGION					
	>gi 2131678 pir S64439					
	hypothetical protein YGR130c -		}			
	yeast (Saccharomyces		1			
	cerevisiae)		1			
	>gi 1323215 gn1 P1D e243523		1	}		
0.38	(Z72915) ORF YGR130c		į	}		
0.20	[Saccharomyces cerevisiae]	1723710	<none></none>	<none></none>		
0.26	(AF046125) immediate early 2			ZNUNES	<none></none>	46
0.20	[Rat cytomegalovirus]	2996117	<none></none>	<none></none>		
0.024	(AF102855) synaptic SAPAP-			ZNUNES	<none></none>	47
-	interacting protein Synamon	4151809	<none></none>	<none></none>		
1	(AF040954) putative protein			CHOINE	<none></none>	48
1	phosphatase I nuclear targeting	1			1	1
0.017	subunit [Rattus norvegicus]			1.	1	1
3e-0	(D90914) hypothetical protein	2773341	<none></none>	<none></none>	-NONTE:	١
1	HYPOTHETICAL 100.6 KD	1653522	<none></none>	<none></none>	<none></none>	49
1	TRP-ASP REPEATS	i		-	<none></none>	50
1	CONTAINING PROTEIN	1	l		1	1
:	C2C6.04C IN CHROMOSOME	1			1	1
3e-0	1 CZCG.04C II. C. II. C. II.				1	}
1	- 1	3219965	<none></none>	<none></none>	<none></none>	ے ا
1	(AF115480) cAMP-dependent	1		_	CHOME	5
1	In an ananine nucleotide	1	1	1	(1
[] 7e-C	exchange factor [Mus musculus	1105517	1		S	1
	CACHAII go 100101	4185567	<none:< td=""><td><none></none></td><td><none:< td=""><td>5</td></none:<></td></none:<>	<none></none>	<none:< td=""><td>5</td></none:<>	5

		The Wheel we Go	nbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
fiji.	Nearest N	eighbor (BlastN vs. Ge	illoanik)			· ·		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION		P VALUE		
<u> </u>	ACCESSION		7		HYPOTHETICAL 43.2 KD			
					PROTEIN C34E10.1 IN	1		
1		!			CHROMOSOME III	ı		
					>gi 500724 (U10402) C34E10.1			
	·	Ī			gene product [Caenorhabditis			
		<none></none>	<none></none>	1176527	elegans)	3e-20		
53	<none></none>	G.pallida repetitive			beta-globin - chimpanzee	0.6		
	¥96444	DNA element	5.0	2118936	(fragment)	8.6		
54	X85444	DIVA Cicinent			MICROTUBULE-			
	1				ASSOCIATED PROTEIN 1A	1		
	.	Synechococcus sp.		<u>'</u>	microtubule-associated protein			
	ł	cpeB, cpeA genes and			MAP1A - rat >gi 205538	2.2		
55	X72961	ORF3	5.0	462569	norvegicus]	2.2		
	A12301	O.Q.S		1	1	ļ		
	1	Human WD repeat				,		
	1	protein HAN11	ł		(Z67990) similar to cuticle	1.3		
56	U94747	mRNA, complete cds	5.0	3875538	collagen	 		
				1	`	•		
	1	Homo sapiens	l	}	\	1		
		integrin alpha-7			collagen - Paralvinella grasslei	0.002		
57	AF032108	mRNA, complete cds	5.0	2147194	ASPARTYL-TRNA			
			l		SYNTHETASE synthetase			
}		G.gallus mRNA for	1	3122885	[Bacillus subtilis]	3e-11		
58	Z50798	p52	5.0	3122803	[Dacinus scottie]			
		Human mRNA for	1	<u>.</u>	(Y15513) Prodos protein	ł		
1	1	KIAA0386 gene,	5.0	2632098	[Drosophila melanogaster]	9e-12		
59	AB002384	complete cds	3.0	2032070				
	t	Thermofilum penden		1	ì	1		
1	1	DNA for 16S and	"]	}	1			
	1	23S ribosomal RNA,	ł		1	1		
1	l	tRNA-Met, and tRNA			}:	**********		
١.,	*******	1	4.9	<none></none>	<none></none>	<none></none>		
60	X 14835	Gly	 		NONSTRUCTURAL	1		
1	1		1		PROTEIN NS-S spotted wilt	- I		
1	1	Hordeum vulgare		1	virus (strain CPNH1) non-			
	1	nucellin gene,	1	j	structural protein [Tomato	2.8		
61	U87149	complete cds	4.9	128578	spotted wilt virus]	 		
-	307149	Mus musculus gene	T		HYPOTHETICAL PROTEIN	1		
1	1	for integrin alpha v	l	Ţ	UL61 cytomegalovirus (strain			
1	ì	subunit, promoter			AD169) cytomegalovirus]	0.038		
6	D87541	region	4.9	136956	AD 1091 Cytomegalovitus)	1-3:22		
<u> </u>	1		1	1	· l	}		
	1	Mus musculus mena		1	(AB007934) KIAA0465 protei	in l		
	}	protein (Mena)		2413002	[Homo sapiens]	6e-07		
ے ا	3 U72520	R -	ls 4.9	3413892	(fuomo sabicus)			

		Marklas Go	abank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
14.	Nearest N	Neighbor (BlastN vs. Ge	noank)	1100000]	
SEC	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
	1						
_		enzymatic				i	
1	1	glycosylation-			!		
1	1	regulating gene [rats.			<u> </u>	i	
١		Sprague-Dawley,			1	1	
		streptozotocin	·			1	
 		diabetic, heart,	4.0	<none></none>	<none></none>	<none></none>	
64	S79797	mRNA, 5010 nt]	4.8	CHOILE	RECEPTOR RECOGNIZING		
				{	PROTEIN gp38 - phage Ox2	- l	
1	1		•		>gi 15126 (X05675) gene 38	•	
1		Homo sapiens mRNA			(AA 1-266); pid:g15126	· 1	
1		for KIAA0530	4.8	138022	[Bacteriophage Ox2]	3.6	
6	AB011102	protein, partial cds Penacus monodon	7.0			į	
		phosphopyruvate		1		ļ	
	1	hydratase mRNA.	Ì	1	(D16221) endochitinase [Oryza		
١,		complete cds	4.8	500615	sativa]	2.8	
6	AF100985	Complete cus			(AL021471) similar to		
	1	Bacillus subtilis		,	Eukaryotic aspartyl proteases	1	
1	- 1	gamma-	ì	ł	[Caenorhabditis elegans]	1	
	j	aminobutyrate	l		Eukaryotic aspartyl proteases	2.8	
6	7 U31756	permease cds	4.8	3880699	[Caenorhabditis elegans]	2.0	
F	-			}		}	
1	1	Pisum sativum	·	l	Į.	<u> </u>	
Ŋ.	1	chloroplast	ļ		1	1	
1	1	processing enzyme				ŀ	
	1	mRNA, nuclear gene	1		(U83658) FH1/FH2 protein	[
- 1	. !	encoding chloroplast	1	1800145	homolog [Emericella nidulans]	1.6	
يا ا	8 U25111	protein, complete cds	. 4.8	1800143	Homolog (S		
	1	Mus musculus Cdx-2			·		
		homeobox protein	4.7	<none></none>	<none></none>	<none></none>	
1	9 U00454	gene, complete cds. Hamster c-Ha-ras	 		RENIN-BINDING PROTEIN	1	
1	1	protein gene,	1	} .	(RNBP) protein [Rattus		
1.	10 1404166	complete cds.	4.7	1710606	norvegicus]	0.88	
-	0 M84166	Mus musculus major		1			
1	1	sperm fibrous sheath		1	1	1	
1	Į	protein Pro-	1	1	1	1	
-		mAKAP82 gene.	1	· ·		ł	
- {		alternative splice	1			<none></none>	
	71 AF087516	1	4.6	<none></none>	<none></none>	SINOINES	
-		M.esculenta mRNA			1	l	
		for granule-bound			NONTE	<none></none>	
- {	72 X74160	starch synthase	4.6	<none></none>	<none></none>	+=	
-		Haloferax volcanii		,	j	j	
1	1	superoxide dismutas	e į		(AC002409) putative ubiquitin		
- 1		(sod2) gene, comple	te	2622207	protease [Arabidopsis thaliana]	3.4	
- 1	73 M97487	cds.	4.6	2623307	Iprotease (Atabidopois C.E.Terror)		

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		ni Nin Go	obank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	Nearest ?	leighbor (BlastN vs. Ge	ndank)	1100001		1	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION 1	VALUE	
-		Drosophila			 		
		melanogaster				NOTE:	
		suppressor of sable	4.5	<none></none>	<none></none>	<none></none>	
74	M57889	gene, complete cds. Rattus norvegicus				l l	
	1	mRNA for RNA				<none></none>	
			4.5	<none></none>	<none></none>	SNOWES	
75	D49708	binding protein					
	1	Yeast GTS1 gene for		}	DETTS		
	1	glycin-threonin/serine			(U42580) NETTF (7x), DETTS		
	1	repeat protein,		1	(4x) [Paramecium bursaria	3.3	
	201052	repeat protein,	4.5	. 2447195	Chlorella virus 1]	3.3	
76	D31853	complete cds Human partial cDNA					
ļ		sequence, clone				<none></none>	
		bs613;	2.9	<none></none>	<none></none>	CNOINE	
77	Z47036	DS013,]	
ł		Rattus norvegicus	ţ	1	j		
Ì	ł	gastric inhibitory	1	Ì			
1	ł	peptide receptor	·		(AF007871) torsinA [Homo	2e-07	
	L19660	mRNA, complete cds	2.7	2358279	sapiens] immediate-early protein E110 -		
78	L19000	IIII CATA, COMPA			immediate-early protein in 12110	1	
1	1		1	7	human herpesvirus I (strain	8.4	
	X82841	A thaliana Aco gene	2.6	483212	HFEM) (fragment)	 	
79	A82841	A.diditale			1	ľ	
1	į.		Į.		1		
1	1	S.purpurascens fam A				1	
1	i i	and famB genes for		1		l	
}	1	FAS domain and acy	너		(U95031) sublingual gland	1	
1		CoA-dehydrogenase:	s.	********	mucin [Homo sapiens]	0.47	
81	X61931	respectively	2.6	2290534	mucin (Home seprens)		
1-8	1	Human lactate	l	\$		}	
	1	dehydrogenase-C	1	1	(AB007874) KIAA0414 [Home	o {	
		(LDH-C) mRNA,		2007440	sapiens]	3.1	
8	1 U13680	complete cds.	2.5	2887449	Judicino		
\ <u></u>		Homo sapiens	1		(AB008859) pheromone		
		KIAA0409 mRNA.	1	3130157_	receptor [Fugu rubripes]	5.4	
8	2 AB00786	9 partial cds	2.4	713017/			
		H.sapiens mas proto	-	<none></none>	<none></none>	<none< td=""></none<>	
8	3 X97479	oncogene. 5' region	2.1_				
	1		. 1	\$	ł –		
	1	R.norvegicus mRN	1.9	<none></none>	<none></none>	<none< td=""></none<>	
	84 X98374	for KIS protein	1.9	1 - 3,0,,5		1	
					}	l l	
1	1	Aquifex aeolicus	e			NO.	
	1	section 42 of 109 o		<none></none>	<none></none>	<none< td=""></none<>	
- 1	85 AE0007	10 the complete genon	1.9				

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			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
Nearest N	leighbor (BlastN vs. Ge	nbank)	Nearest Neight				
ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
	Homo sapiens mRNA						
	for repressor protein,	1.9	<none></none>	<none></none>	<none></none>		
D30612	Homo saniens						
			1	NONE	<none></none>		
V1/221		1.9	<none></none>	<nunes< td=""><td>3,10</td></nunes<>	3,10		
114321	E coli genomic DNA,			(DZ0205) DNA hinding protein	1		
}	Kohara clone	i			7.9		
D90773	#262(30.3-30.5 min.)	1.9	1536816	[Chlorella virus]			
B30713	Archaeoglobus		1	(Y70095)	}		
1	fulgidus section 116	1		oursyste orthophosphate	1 .		
<u>.</u>	of 172 of the	_	500645	dikinase (Flaveria trinervia)	2.7		
AE000991	complete genome	1.9	320043	GIRTIMSC IS 12.			
	Rattus norvegicus	1			1		
1	p95 Vav (Vav) proto-	· i	}	(AL023496) hypothetical	1		
1	oncogene mRNA.	1	4159178	erotain	1.6		
U39476	complete cds.	1.9	4136176	HYPOTHETICAL PROLINE-	1		
1		l l	ì	RICH PROTEIN KIAA0269	1		
1	1	ì		>gi 1665805 gnl PID d1014089	1		
1		1		(D87459) Similar to Volbox	1		
1	Human transcription	.1	i .				
1	. i	4 10	2495730	[Homo sapiens]	0.23		
U28838	subunit				1		
Į.	Darbie porvegicus	1	1				
1	Rattus not regions			UL47h protein - Marek's diseas	0.23		
7/20106	mpNA complete cd	s. 1.9	478380	virus	- U.23		
0.20100	mikiva, complete c				i i		
ł	Mouse mammary	1		(A COOADIO) similar to Leucine	<u>.</u> [
1	tumor virus putative	·	Į.	Lish masmembrane proteins:	1		
· ·		į		144% similarity to 1142767	1		
1	polyprotein, and	- (·	(DID: 21736918) [Homo			
1	superantigen mRNA	. ,			4e-33		
AF07101		1.9	2/81386	Sapiens			
		1	·	1	}		
			1	1	1		
1	c-fos proto-oncogei	ne j		1			
ł		1,	<none></none>	<none></none>	<none< td=""></none<>		
4 AF06188	l complete cds	- 1.8	1 3101.55		1		
	Plasmodium		1 .				
1	falciparum	1	1				
L L				•			
	chromosome 2.	İ	į	·	1		
	section 34 of 73 of the complete			<none></none>	<none< td=""></none<>		
	D30612 Y14321 D90773 AE000991 U39476 U28838 U20106	ACCESSION DESCRIPTION Homo sapiens mRNA for repressor protein, partial cds Homo sapiens PMP69 gene, exons PMP69 gene, exons Second PMP69 gene, exons Y14321 8,9.10 & 11 E.coli genomic DNA, Kohara clone #262(30.3-30.5 min.) Archaeoglobus fulgidus section 116 of 172 of the complete genome Rattus norvegicus p95 Vav (Vav) proto- oncogene mRNA, complete cds. Human transcription factor TFIIIB 90 kD subunit Rattus norvegicus synaptotagmin VII mRNA, complete cd Mouse mammary tumor virus putative integrase, env polyprotein, and superantigen mRNA complete cds Mesocricetus auratu c-fos proto-oncoger protein (c-fos) gene A F061881 complete cds	ACCESSION DESCRIPTION P VALUE Homo sapiens mRNA for repressor protein, partial cds PMP69 gene, exons Y14321 8.9.10 & 11 E.coli genomic DNA, Kohara clone D90773 #262(30.3-30.5 min.) Archaeoglobus fulgidus section 116 of 172 of the AE000991 complete genome Rattus norvegicus p95 Vav (Vav) proto- oncogene mRNA. complete cds. 1.9 Human transcription factor TFIIIB 90 kDa subunit U20106 mRNA. complete cds. 1.9 Mouse mammary tumor virus putative integrase, env polyprotein, and superantigen mRNA, complete cds 1.9 Mesocricetus auratus c-fos proto-oncogene protein (c-fos) gene, complete cds 1.8	ACCESSION DESCRIPTION P VALUE ACCESSION Homo sapiens mRNA for repressor protein, D30612 partial cds Homo sapiens PMF69 gene, exons PMF69 gene, exons Y14321 8,9.10 & 11 1.9 <none> E.coli genomic DNA, Kohara clone D90773 #262(30.3-30.5 min.) 1.9 1536816 Archaeoglobus fulgidus section 116 of 172 of the complete genome Rattus norvegicus p95 Vav (Vav) proto- oncogene mRNA. U39476 complete cds. 1.9 4158178 Human transcription factor TFI11B 90 kDa subunit U23838 subunit U20106 mRNA, complete cds. 1.9 478380 Mouse mammary tumor virus putative integrase, env polyprotein, and superantigen mRNA, complete cds 1.9 2781386 Mesocricetus auratus c-fos proto-oncogene protein (c-fos) gene. complete cds 1.8 <none></none></none>	ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION		

	Nanzari	Neighbor (BlastN vs. G	enbank)	Nearest Neighbo	r (BlastX vs. Non-Redundant Prot	
SEQ			P VALUE	ACCESSION	DESCRIPTION	P VALUE
ID	ACCESSION	DESCRIPTION	PVALUE			
		Horseshoe crab	<u> </u>			
		mRNA for	1			
	1	coagulation factor B,	1	<none></none>	<none></none>	<none></none>
96	D14701	complete cds	1.8	2140112		
		P.falciparum	ì			
	1	multidrug resistance	1		1	
	1	(MDR) gene,	1	<none></none>	<none></none>	<none></none>
97	M29154	complete cds.	1.8	- CINOTIES		
<u> </u>	1	Rattus norvegicus		•		
	1	(clone pCNPII) 2',3'-	· {	1	1	
	1	cyclic nucleotide 3'-	. }	1 .	,	
l	1	phosphodiesterase	1 .	1		1
!	1	(CNPII) mRNA,	1	<none></none>	<none></none>	<none></none>
98	L16532	complete cds.	1.8	KNONL		l
	1	Plasmodium	1	1		1
1	1	falciparum	1		1	
1	1	chromosome 2,	1	1		
}	1	section 71 of 73 of	l l			
1		the complete		<none></none>	<none></none>	<none:< td=""></none:<>
99	AE00143	4 sequence	1.8	CHOILE		1
1-		D.melanogaster ger	ie			
1	1	for protamine	١.,	<none></none>	<none></none>	<none:< td=""></none:<>
10	o Z46785	(mst35Bb).	1.8	- CHOINE		1
**		P.sylvestris mRNA	• 1	· I	(D90452) 1-caldesmon I [Homo)]
1	1	for glutamine	1.8	219896	(iama)	9.7
1 10	1 X6982	synthetase	1.0		INSULIN-LIKE GROWTH	J.
			· \	ł ·	FACTOR BINDING PROTEIL	`\
1	l l		1		4 (IGFBP-4) (IBP-4) (IGF-	
1	1	}	1		BINDING PROTEIN 4) factor	-
	1	1	j	i i	binding protein-4 - sheep	-
1	ł		1		(fragment) factor-binding	1
l	I	Rattus norvegicus	.,,_ {	1	protein-4, IGFBP-4 (sheep,	
1	ļ	CTD-binding SR-	like		liver, Peptide, 237 aa] [Ovis	2.5
1.	1.	protein rA8 mRN	A, 1.8	2497252	aries]	
1	02 U490.	5 complete cds	1.8			1
		Homo sapiens		l l		1
ļ	1	kallistatin (PI4) g	ere.		(AC005223) 55585	2.4
1	1	exons 1-4, compl	ete 1.8_	4204267	[Arabidopsis thaliana]	
	103 L281	01 cds				1
		Pandorina morun	,	}		.]
1	l l	internal transcrib	ed			- 1
1	l l	internal transcrib		1	}	1
1		spacer 1, 5.8S	anne	1	1	1
1	l	ribosomal RNA	gene.	1		1
l	l l	and internal	,		(Z99121) permease [Bacillus	1.9
- {	1	transcribed space	er 2.	2635909	subtilis)	
- 1	104 U66	987 complete sequer	1.0			

PCT/US00/18374 WO 01/02568

	Niconae Ni	eighbor (BlastN vs. Ge	nbank)	Nearest Neight	oor (BlastX vs. Non-Redundant Pro	Citis)
# 10 pm	Nearest IV	eignbor (Blaset vs. Co				
EQ			P VALUE	ACCESSION	DESCRIPTION	P VALUE
מו	ACCESSION	DESCRIPTION	PVALUE	ACCESSION		
		Human polymorphic			keratin KAP5.5 - sheep	
		Mspl site DNA		012/070	(fragment) >gi 313722	0.65
105	X58033	(D3S3 locus)	1.8	2136878	(Hagment) - g.jo	
103	750055				(AC004877) sco-spondin-mucin-	
		Human p82 (ST5)		•	like; similar to P98167 uncertain	
	\	mRNA, alternatively		2.00007	[Homo sapiens]	0.64
106	บ15780	spliced, complete cds	1.8	3638957	(U00690) calcium channel alpha	
100	013700	Homo sapiens			1 subunit [Drosophila	
	l	synaptotagmin VII	}		melanogaster]	0.51
	AF038535	mRNA, partial cds	1.8	457927	meianogasteri	
107	AF030333	Homo sapiens clone		l	HOMEOBOX PROTEIN HOX-	
	1	23585 mRNA				0.28
	A TO50124	sequence	1.8	232263	D1 (HOX-4.9)	l
108	AF052134	H.sapiens HEK2			GROWTH-ARREST-SPECIFIC	
	1	mRNA for protein	l	ł	GROW IN-ARREST-SI ECUTE	
	1	tyrosine kinase	1		PROTEIN 1 gene product	0.22
			1.8	1730198	[Homo sapiens]	1
109	X75208	receptor. Xenopus laevis	1		TO THE TONIES CTOP	}
	1	mRNA for SOX-D,		1	TRANSCRIPTION FACTOR	0.17
			1.8	2494501	FKH-4 factor [Mus musculus]	+
110	AB013896	Human HepG2 3'	-		TOTAL A DAE A protein	.
		Human richos	}		(AB007923) KIAA0454 protein	0.002
		region cDNA, clone	1.8	3413870	[Homo sapiens]	0.002
11	D16947	hmd6b10		+		1
	1	Toods	,	}	(AL031174) hypothetical	5e-08
1		Mouse DNA, Tearly	1.8	3393018	protein	36-00
11	2 D13547	alpha (TEA) region			HYPOTHETICAL II.3 KD	}
[ł		PROTEIN C2C6.07 IN	
1	1		,	1	CHROMOSOME I	
1	1		1	1	>gi 2370504 gnl P1D e339194	1
l	i		}	1	pombe]	. 1
1	1		Ţ		>gi 345 1305 gnl PID e1316730	' }
1	l	l l	1	1	(AL031324) very hypothetical	1
1	- 1			1	protein [Schizosaccharomyces	.
1	l .	Woodchuck c-myc		3183405	pombel	8e-10
1	13 M35498	protein gene, exon	1. 1.8	7103403		1
		Hamster c-Ha-ras	1	Į.	(AC004665) unknown protein	
1	1	protein gene.	١.,	3386622	[Arabidopsis thaliana]	2e-10
1	14 M84160	complete cds.	1.8	3300022		1
 -		Mychodea carnosa	. 1		1	1
1	1	18S ribosomal RN	A.		(AC005306) R27216_1 [Hom	0
1	ł	gene, complete		1224002	[sapiens]	3e-2
١,	15 U3313	sequence	1.8	3334982		
-		Homo sapiens		1		1
-	1	putative tumor	ļ	1		- [
1		suppressor (BIN1)			<none></none>	<non< td=""></non<>
1	16 U8400		1.7	<none></none>	CHOILE	

		(D) N (C)	nbank) T	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
. Viv.	Nearest N	leighbor (BlastN vs. Ge	nbank)				
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Borrelia burgdorferi					
		(section 7 of 70) of				NONT	
117	AE001121	the complete genome	1.7	<none></none>	<none></none>	<none></none>	
117	712001121	Archaeoglobus				İ	
		fulgidus section 165		<u> </u>		Į.	
		of 172 of the	į			ALONTES	
118	AE001114	complete genome	1.7	<none></none>	<none></none>	<none></none>	
		Angiostrongylus				1	
	l	cantonensis adult-				1	
	•	specific muscle					
		protein-1 gene. partial			<none></none>	<none></none>	
119	U82064	cds	1.7	<none></none>	<nunes< td=""><td>CHOILE</td></nunes<>	CHOILE	
		Buchnera aphidicola				}	
	1	plasmid pLeu-Sg.					
	1	complete plasmid	_		<none></none>	<none></none>	
120	AF041836	sequence	1.7	<none></none>	ANOINE	- 1	
		Lymnaea stagnalis		ļ		.	
	Ì	FMRFamide gene.		MONTES	<none></none>	<none></none>	
121	M87479	mature peptides.	1.7	<none></none>	<u> </u>		
		,]			j	
		Xenopus laevis	·			1	
	i	fibroblast growth	ļ	`		1	
	1	factor receptor	1.7	<none></none>	<none></none>	<none:< td=""></none:<>	
122	M55163	mRNA, complete cds.	1./				
1	Ì		ł	1			
١.,,	057565	receptor [rats, Genomic, 1928 nt]	1.7	<none></none>	<none></none>	<none:< td=""></none:<>	
123	S57565	Simian	1 - 11			1	
Į		immunodeficiency				İ	
	1	virus (SIV) pol		1			
124	M27256	region.	1.7	<nońe></nońe>	<none></none>	<none:< td=""></none:<>	
129	14127230	Human chromosome				İ	
] .		8 anonymous clone	1			<none:< td=""></none:<>	
125	U31516	pBS8-165	1.7	<none></none>	<none></none>	ZNONE.	
 ~~	1	Human gene for			j		
1		heterogeneous		1	1	1	
ľ		nuclear	į.	•	1	1	
1	1	ribonucleoprotein	1	ł	Ì		
	1	(hnRNP) core proteir	1		NONTE	<none< td=""></none<>	
120	5 X12671	AI	1.7	<none></none>	<none></none>	SHORE	
		Paeonia suffruticosa	1 .	1		1	
1	l	ssp. spontanea		S	}	1	
1	1	alcohol	1	1		}	
1	1	dehydrogenase iB	.1	1		1	
	1	(AdhlB) gene, partia		NONE	· <none></none>	<none< td=""></none<>	
12	7 AF00905	cds	1.7	<none></none>	710112		

				Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
7.100 7.000 7.000	Nearest N	leighbor (BlastN vs. Ge	nbank)	Nearest Neighbo	I (Blaste vs. 1.c.		
SEQ		DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	PVALUE	
Œ	ACCESSION	DESCRIPTION					
		Mus musculus				1 . 1	
		transketolase gene,				<none></none>	
		transketotuse gene,	1.7	<none></none>	<none></none>	SNOINES	
128	AF046917	exon 6 and partial cds Homo sapiens mRNA				- 1 1	
	l					1	
	į.	for Acyl-CoA				NONTE	
		synthetase 3,	1.7	<none></none>	<none></none>	<none></none>	
129	D89053	complete cds Staphylothermus				1 1	
		Staphylothermus			į (-	
	{	marinus surface layer-	1		į	1 . 1	
	1	associated STABLE	}	1 , .	·		
		protease gene,	1.7	<none></none>	<none></none>	<none></none>	
130	U57968	complete cds.	1./	<u> </u>		iii	
		Bovine herpesvirus 1	}	-		1	
1	1	(clone p95) UL24	1				
1	1	homologue gene.	1.7	<none></none>	<none></none>	<none></none>	
131	L39072	complete cds.	1./	1 (110.12		ì	
		Drosophila simulans	1.	i	\		
1		retrotransposon 297				1	
}	1	5'-LTR and flanks	1.7	<none></none>	<none></none>	<none></none>	
132	X04980	(pWK1020)	1./			1	
		Archaeoglobus		1	1	 	
l		fulgidus section 165	1	1			
	1	of 172 of the	1.7	<none></none>	<none></none>	<none></none>	
133	3 AE001114	complete genome	1./_				
	,	Human mRNA for	ì	1			
1	t	insulin-like growth	1.7	<none></none>	<none></none>	<none></none>	
13	4 X04434	factor 1 receptor	1./			•	
		Mus musculus	. 1	ŀ	1	İ	
1		C57BL/6J epiderma	,)	i .	· ·	1	
ł		surface antigen	}	! .			
1	1	(mesa) mRNA.	1.7	<none></none>	<none></none>	<none></none>	
13	5 U07890	complete cds.	1./	- 110112		1	
		Human tyrosinase	(ł]	
]	gene. 5'-flanking	1	I			
1	1	region cell-specific	1.7	<none></none>	<none></none>	<none:< td=""></none:<>	
13	36 D26163	transcription)				į į	
		Panorpa nipponens	³				
- 1	l l	NADH	•1		1	1	
	į	dehydrogenase	1	}	1	1	
- [1	subunit 5 gene.	1	1	1		
į.	Į	mitochondrial gene	1		1		
-		encoding	1	1	İ	!	
- }		mitochondrial		<none></none>	<none></none>	<none< td=""></none<>	
- 1 .	37 AF0938	8 protein, partial cds	1.7	_1		_	

PCT/US00/18374

te la	Negreet N	leighbor (BlastN vs. Ge	nbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
	METICSLI	reignbor (Blast 13: Co						
SEQ	_	TERCH WILLIAM	DVALUE	ACCESSION	DESCRIPTION	P VALUE		
ID	ACCESSION		P VALUE	ACCESSION				
		Xenopus laevis						
		mRNA for			;			
		cytochrome P-450,		NONE	<none></none>	<none></none>		
138	D50560	complete cds	1.7	<none></none>	CHORES	4101102		
		Mus musculus			}			
1		phospholipase D1	i		ł i			
		(PLD1) gene, exons						
	1	18 and 19, complete		NONT:	<none></none>	<none></none>		
139	AF083488	sequence	1.7	<none></none>	CNOILE			
		Mus musculus						
		Pontin52 mRNA,		NONES	<none></none>	<none></none>		
140	AF100694	complete cds	1.7	<none></none>	40112			
		C+						
	į l	Streptococcus			}			
	1	salivarius	· ·	·				
	į	thermophilus beta-D-						
	1	galactose (lacZ) gene.		į				
		complete cds. > ::		ł	1			
		gb[M63636]STRLAC		İ	i			
	}	ZZ Streptococcus	1	1		[
	1	thermophilus beta-D-				1		
	1	galactosidase (lacZ)	1	<none></none>	<none></none>	<none:< td=""></none:<>		
141	M73749	gene, complete cds.	1.7	KNOINES				
	1	Archaeoglobus		1	Ì			
	1	fulgidus section 165	l	1	(U84971) unknown [Homo	1		
	I	of 172 of the		2183023	sapiens]	9.2		
142	AE001114	complete genome	1.7	2183023	GENOME POLYPROTEIN			
	Į.		1		CONTAINS: N-TERMINAL	1		
	i]		PROTEIN (P1); HELPER	ł		
			l .		COMPONENT PROTEINASE	1		
	}				INCLUSION PROTEIN (CI); 6	1		
	1		{ ·	Į ,	KD PROTEIN 2 (6K2);	}		
!	}	Thuman men 137	ŀ		GENOME-LINKED PROTEIN	į		
	1 .	Human type IV	}	1	(VPG); NUCLEAR virus	1		
		sodium channel alpha	1.7	130504	(strain D)	9.2		
143	L01983	polypeptide Plecotus rafinesquii	1./	150504		1		
		mitochondrial	1	1		ł		
}	1		.1	l	(AB014541) KIAA0641 protein	1		
		cytochrome b gene. 5	1.7	3327096	[Homo sapiens]	9.1		
144	L19731	end. Archaeoglobus	 *-'	332.030	(
1	1	fulgidus section 165	1	-		1		
1	1	of 172 of the	1	1	(U84971) unknown [Homo]		
١		1	1.7	2183023	sapiens	8.8		
145	AE001114	complete genome	<u> </u>					

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- F#S70	Nearest N	leighbor (BlastN vs. Go	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	14emest 1	reigitooi (Diastra va. Or					
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
			_				
	·	Bos taurus serum amine oxidase mRNA, complete cds.			SIGNAL TRANSDUCER AND		
		> oxidase=amiloride- binding protein			ACTIVATOR OF TRANSCRIPTION 6 (IL-4 STAT) >gi 559855 (U16031) IL-	· •	
146	L27218	homolog [cattle, liver, mRNA, 2664 nt]	1.7	1174459	4 Stat [Homo sapiens]	7.1	
		Caenorhabditis elegans cosmid W07E11, complete sequence			(4.0005033) 10400		
147	Z49868	[Caenorhabditis elegans]	1.7	4204263	(AC005223) 40409 [Arabidopsis thaliana]	6.7	
	247000	·		*	PERIPLASMIC NITRATE REDUCTASE PRECURSOR >gi 1086107 pir S50163 nitrate		
		Caenorhabditis elegans cosmid F32F2, complete sequence [Caenorhabditis			reductase large chain precursor, periplasmic - Thiosphaera pantotropha >gi 600093 (Z36773) periplasmic nitrate reductase large subunit		
148	AL022271	elegans]	1.7	2497969	[Paracoccus denitrificans]	6.7	
149	U43844	Mus musculus cyclin D3 gene, complete cds	1.7	3861490	(AF062037) capsid protein precursor [Thosea asigna virus]	5.1	
150	Z25464	S.cerevisiae UNF1, LTV1, MRP8, CYB3 and TGL1 genes, complete CDS's	1.7	1255404	(U53151) weak similarity to cytochrome b [Caenorhabditis elegans]	4.1	
151	U77846	Human elastin gene, partial cds and partial 3'UTR	1.7	3355682	(AL031124) putative secreted lyase	4.0_	
152	X62880	S scrofa mRNA for calcium release channel (CRC)	1.7	3327080	(AB014533) KIAA0633 protein [Homo sapiens]	4.0	
153	Y00067	Human gene for neurofilament subunit M (NF-M)	1.7	. 479829	heterogeneous ribonuclear particel protein homolog - Caenorhabditis elegans similarity to RNA recognition motifs [Caenorhabditis elegans]	3.9	

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75.25E	Nearest N	leighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
- V. A. A. A.	r con cat i	CIENOU (DIESE: 13. O.					
SEQ		DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
B	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	T DESCRIPTION	1 VALUE	
					(AC000106) Contains similarity		
1 1			·		to Rattus AMP-activated protein		
1		D.melanogaster gene	1		kinase (gb X95577).		
1 1		for Beta-tubulin,		2240782		3.8	
154	X68393	exons 1 and 2	1.7	2342682	[Arabidopsis thaliana]	3.0	
1 1			1		(ORFIA/IB) [CONTAINS:		
1		GI II	ļ .		RNA-DIRECTED RNA		
1 1		Shuttle vector	1		POLYMERASE; HELICASE;		
		pAUR123 gene for		417704	PROTEASE 1	3.8	
155	AB012284	Aur.1-C, complete cds	1.7	41//04	[FROTEASE]	3.0	
		Rattus norvegicus	l				
1		mitochondrial	i	• •	}		
1 1		intermediate				ł	
1			•		(AE000613) H. pylori predicted		
	100000	peptidase (MIP)	1.7	2314209	coding region HP1054	3.1	
156	M96633	mRNA, complete cds.	1.7	2314209	INSULIN-LIKE GROWTH		
)			1		FACTOR BINDING PROTEIN	ì	
ł			ij		4 (IGFBP-4) (IBP-4) (IGF-	ł	
1			1 1		BINDING PROTEIN 4) factor-	1	
]					binding protein-4 - sheep		
1		Rattus norvegicus			(fragment) factor-binding		
		CTD-binding SR-like	ì [protein-4, IGFBP-4 [sheep,		
i		protein rA8 mRNA,			liver, Peptide, 237 aa] [Ovis	ì	
157	U49055	complete cds	1.7	2497252	aries]	3.0	
13/		complete eds					
1	l	Mus musculus mRNA	}		iduronate-2-sulfatase, IDS (EC		
1		for myc-intron-	1		3.1.6.13) Peptide Mutant, 550		
158	Y15907	binding protein-1	1.7	912776	aal	3.0	
T-133		Methanococcus					
1	i e	jannaschii section 142					
1		of 150 of the			(AF052252) fork head domain		
159	U67600	complete genome	1.7	2982355	protein FKD9 [Danio rerio]	3.0	
					·		
}		Homo sapiens				}	
1	1	calumein (Calu)	i I		(AF052252) fork head domain		
160	AF013759	mRNA, complete cds	1.7	2982355	protein FKD9 [Danio rerio]	2.9	

illight y	Nearest N	leighbor (BlastN vs. Go	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ						1	
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					Human niRNA product		
					KIAA0077 (TR:Q14997);		
}					cDNA EST yk243h8.5 comes		
l l					from this gene; cDNA EST]	
{					yk243h8.3 comes from this		
1					gene; cDNA EST yk359h4.5		
()					comes from this gene	1	
}					[Caenorhabditis elegans]	<u>[</u>	
]					>gi 3880318 gnl PID e1349839	{	
1					(Z81133) Similarity to Human	1	
į į					mRNA product KIAA0077	{	
i '	·			,	(TR:Q14997); cDNA EST	1	
					yk243h8.5 comes from this		
1	1	Arabidopsis thaliana			gene; cDNA EST yk243h8.3		
1	1	putative transcription			comes from this gene; cDNA		
		factor (MYB90)			EST yk359h4.5 comes from this		
161	AF062915	mRNA. complete cds	1.7	3878065	gene	2.3	
		H.sapiens genomic	1		(AC004877) sco-spondin-mucin- like; similar to P98167 uncertain		
		DNA (chromosome	1.7	3638957	[Homo sapiens]	2.3	
162	X87526	3; clone NL3003R)	1./	3038937	[Hollio Sapiens]	2.5	
1		Homo sapiens			(AF005632) phosphodiesterase	1	
1		chromosome 5, PAC			Unucleotide pyrophosphatase	i .	
163	AC005573	clone 202e13	1.7	2465540	beta [Homo sapiens]	1.8	
		Homo sapiens gene				1	
1		for prostacyclin					
1	_	synthase, exon 10 and			steroid hormone receptor TR3 -	, ,	
164	D83402	complete cds	1.7	627608	human sapiens}	1.7	
	1	Homo sapiens deltex			(AB007864) KIAA0404 [Homo		
		(Dx) mRNA,	, ,	2662089	(AB00/864) KIAA0404 (Hollio sapiens)	1.7	
165	AF053700	complete cds	1.7	2002089	Sahieral	 	
		Mus musculus 6-		,	1		
ł	·	pyruvoyl-			1	1	
1	1	tetrahydropterin			(AF006564) alcohol	1	
-		synthase (Pts)			dehydrogenase [Drosophila		
166	AF043225	mRNA, complete cds	1.7	2352538	persimilis) persimilis)	1.4	

	Nearest	Neighbor (BlastN vs. G	enhank)	Negreet Noigh	ibor (BlastX vs. Non-Redundant Pr	mtoine)
SEO		Vergiloof (Diase) vs. C	l libank)	14cmear 14cial	Tool (Blaster Vs. 14011-Redundant P	Toteins)
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUI
		Thornis aquatique				I
		Thermus aquaticus thermophilus NADH				
		dehydrogenase I				
1		subunits NQO7		'	1	j
1 1		NQ06, NQ05,			1	1
		NQ04, NQ02,			ŀ	}
		NOO1, NOO3,			\$	Ī
		NQ08, NQ09,			ì	
		NQ010, NQ011,			1	}
		NQO12, NQO13, and	l l		(AB006631) The human	
	-	NQO14, complete	1	• .	homolog of mouse Cux-2	1
167	U52917	cds.	1.7	2564334	[Homo sapiens] (2/3425) Similarity to Yeast	1.0
					(2/3425) Similarity to Yeast	
' <u> </u>					hypothetical YIK9 protein	
l					(SW:YIK9_YEAST); cDNA	
!			Į.		EST EMBL:T01252 comes from this gene; cDNA EST	
1			j	•	EMBL:D33205 comes from this	
- 1			1		gene; cDNA EST	
ĺ			i		EMBL:D33955 comes from this	
		M.musculus gene for			gene; cDNA EST	
168	X72222	serotonin 2 receptor	1.7	3875796	EMBL:D35484 co	1.0
					(X83413) DR5 [Human	
i		Crotalus scutulatus	i		herpesvirus 6] >gi 853972	
169		PLA2-like		052001	(X83413) DR5 [Human	
109	U23186	pseudogene	1.7	853971	herpesvirus 6]	0.99
1		Mus musculus factor	ľ		}	
i	. 1	VIII-associated	į.			
	· ·	protein (f8a) mRNA,	- 1		(AC004669) hypothetical	
170	M83118	complete cds.	1.7	3201617	protein [Arabidopsis thaliana]	0.80
T					(AL031282) dJ283E3.3.2 (Céll	
- 1	-	Ì	l	•	Division Cycle 2-Like 2	•
1	1.		- 1		(PITSLRE, p58/GTA,	1
.	1	E.coli ATP-			Galactosyltransferase	į
		dependent proteinase (lon) gene, complete	}		Associated Protein Kinase))	
171		cds.	1.7	4140322	(isoform beta 2-2) [Homo	
			1.1	4140322	sapiens] HYPOTHETICAL PROLINE-	0.78
		1	1		RICH PROTEIN KIAA0269	ł
	Ì		}		>gi 1665805 gnl PID d1014089	1
- 1	1	Human transcription			(D87459) Similar to Volbox	į
1	1	actor TFIIIB 90 kDa	ł		carteri extensin (S22697)	ł
172	U28838 s	ubunit	1.7	2495730	(Homo sapiens)	0.62

-1/2°	Nearest N	Neighbor (BlastN vs. G	enbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant Pr	oteins)
SEO		, , , , , , , , , , , , , , , , , , ,				
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
173	U72487	Rattus norvegicus calcium-independent alpha-latrotoxin receptor mRNA, complete cds	1.7	544411	GLYCOPROTEIN GP100 PRECURSOR (P29F8) discoideum]	0.35
		Aquifex aeolicus section 50 of 109 of	-		FIBROBLAST GROWTH FACTOR RECEPTOR 3 PRECURSOR (FGFR-3) (HEPARIN-BINDING GROWTH FACTOR RECEPTOR) >gi 2117851 pir 155363 fibroblast growth factor receptor 3 - mouse >gi 199145 (M81342) fibroblast growth factor receptor	·
174	AE000718	the complete genome	1.7	2497569	3 [Mus musculus] MACROPHAGE COLONY	0.34
175	AF016897	Oryza sativa GDP dissociation inhibitor protein OsGDI2 (OsGDI2) mRNA, complete cds	1.7	125362	STIMULATING FACTOR I RECEPTOR PRECURSOR (CSF-1-R) (FMS PROTO- ONCOGENE) (C-FMS) factor I receptor - cat >gi 163855 (J03149) M-CSF receptor [Felis domesticus]	0.34
176	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1.7	85058	muscarinic acetylcholine receptor - fruit fly acetylcholine receptor (Drosophila melanogaster) ACROSOMAL PROTEIN SP- 10 PRECURSOR SP-10 - western baboon	0.20
177		Chlamydomonas reinhardtii myosin heavy chain Caenorhabditis elegans cosmid F53B8, complete sequence	1.7	728901	>gi 298488 bbs 127113 (S56458) SP-10=intraacrosomal protein [Papio papio=baboons, Peptide, 285 aa] [Papio hamadryas] (U23517) D1022.7	0.20
178	· · · · · · · · · · · · · · · · · · ·	[Caenorhabditis elegans]	1.7	746516	[Caenorhabditis elegans] >gi 3258651 elegans]	0.068

SEQ D ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION P ACCESSION DESCRIPTION DESCRIPTION P ACCESSION DESCRIPTION DESCRIPTION P ACCESSION DESCRIPTION DE				-ba-le)	Nearest Neigh	bor (BlastX vs. Non-Redundant Pro	oteins)
DESCRIPTION PVALUE ACCESSION DESCRIPTION PVALUE ACCESSION (X87883) mitochondrial capsule sclenoprotein (Ratrus norvegicus) > [il] 154.135 (U48702) mitochondrial capsule sclenoprotein (Ratrus norvegicus) > [il] 154.135 (U48702) mitochondrial capsule (norB) gene. (norB) gene. (norB) gene. (norB) gene sis expressed ubiquitously:; The KIAA0191 gene is expressed ubiquitously:; The KIAA0191 protein trains the C2H2 zince-finger at its N-posphatase norvegicus proponin T isoforms. (Part of the KIAA0191 protein retains the C2H2 zince-finger at its N-posphatase norvegicus proponin T isoforms. (Part of the KIAA0191 protein retains the C2H2 zince-finger at its N-posphatase norvegicus proponin T isoforms. (Part of the KIAA0191 protein retains the C2H2 zince-finger at its N-posphatase norvegicus proponin T isoforms. (Part of the KIAA0191 protein retains the C2H2 zince-finger at its N-posphatase norvegicus proponin T isoforms. (Part of the KIAA0191 protein retains the C2H2 zince-finger at its N-posphatase norvegicus proponin T isoforms. (Part of the KIAA0191 protein retains the C2H2 zince-finger at its N-posphatase norvegicus protein (Part of the KIAA0191 protein retains the C2H2 zince-finger at its N-posphatase norvegicus protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein protein protein (Part of the KIAA0191 protein protein (Part of the KIAA0191 protein pr	av is s Prépas	Nearest N	leighbor (Blastin vs. Ge	noank)	(Acatest recigii		
Raistonia eutropha megaplasmid pHG1 nitric oxide reductase (norB) gene. 179 AF002217 complete cds 1.7 1143538 MCP Rat mRNA for protein tyrosine phosphatase 1.8 D30749 phosphatase 1.7 1228035 terminal region. [Homo sapiens] Rat fast skeletal TnT gene encoding troponin T isoforms. 181 M15202 complete cds. 1.7 T31172 XP2 PRECURSOR Human peroxisome proliferator activated receptor mRNA. 182 L07592 complete cds. 1.7 A033414 SUBUNIT 183 U64031 complete cds. 1.7 A122885 [Bacillus subtilis] Homo sapiens docking protein (DOK-2) mRNA. 184 AF034970 complete cds. 1.7 2289097 [Cricetulus griseus] (AF023270) probable		ACCESSION	DESCRIPTION	P VALUE	ACCESSION		P VALUE
megaplasmid pHG1 nitric oxide reductase (norB) gene. 179 AF002217 complete cds 1.7 1143538 SMCP Rat mRNA for protein tyrosine phosphatase 1.7 1228035 terminal region. [Homo sapiens] Rat fast skeletal TnT gene encoding troponin T isoforms. 181 M15202 complete cds. 1.7 731172 XP2 PRECURSOR Human peroxisome proliferator activated receptor mRNA, 182 L07592 complete cds. 1.7 4033414 4 SUBUNIT ASPARTYL-TRNA SYNTHETASE synthetase 183 U64031 complete cds 1.7 3122885 [Bacillus subtilis] Homo sapiens docking protein (DOK-2) mRNA, complete cds 1.7 2289097 [Cricetulus griseus] Longiflorum mRNA encoding calmodulin. >::: gb[L18912]LILCALM ODU Lilium longiflorum mRNA encoding calmodulin. >::: gb[L18912]LILCALM ODU Lilium longiflorum mRNA encoding calmodulin. >::: gb[L18912]LILCALM ODU Lilium longiflorum mRNA encoding calmodulin.		ACCESSION		T			
AF002217 complete cds 1.7 1143538 SMCP Rat mRNA for protein tyrosine phosphatase 1.7 1228035 terminal region. [Homo sapiens] Rat fast skeletal TnT gene encoding troponin T isoforms. complete cds. 1.7 731172 SKIN SECRETORY PROTEIN XP2 PRECURSOR Human peroxisome proliferator activated receptor mRNA, complete cds. 1.7 4033414 SUBUNIT Dendrobium crumenatum ACC synthase gene. complete cds 1.7 3122885 [Bacillus subtilis] Homo sapiens docking protein (DOK-2) mRNA, complete cds 1.7 2289097 [Cricetulus griseus] 184 AF034970 Lingtiforum mRNA encoding calmodulin. > :: gbjL18912 LILCALM ODU Lilium longiforum mRNA encoding calmodulin mRNA encoding forum mRNA encoding calmodulin. (AF023270) probable			megaplasmid pHG1 nitric oxide reductase			norvegicus] >gi 1354135 (U48702) mitochondia associated cysteine-rich protein	0.039
Rat mRNA for protein tyrosine phosphatase 1.7 1228035 terminal region. [Homo sapiens] Rat fast skeletal TnT gene encoding troponin T isoforms. complete cds. 1.7 731172 SKIN SECRETORY PROTEIN XP2 PRECURSOR Human peroxisome proliferator activated receptor mRNA, complete cds. 1.7 4033414 SUBUNIT LO7592 complete cds. 1.7 4033414 A SUBUNIT Dendrobium crumenatum ACC synthase gene. complete cds 1.7 3122885 [Bacillus subtilis] Homo sapiens docking protein (DOK-2) mRNA, complete cds 1.7 2289097 [Cricetulus griseus] LIO7592 complete cds 1.7 2289097 [Cricetulus griseus]	179	AF002217		1.7	1143538	SMCP	0.039
Rat fast skeletal TnT gene encoding troponin T isoforms. 181 M15202 complete cds. Human peroxisome proliferator activated receptor mRNA, complete cds. 1.7 4033414 PUTATIVE IMPORTIN BETA- 4 SUBUNIT Dendrobium crumenatum ACC synthase gene. 183 U64031 complete cds 1.7 3122885 [Bacillus subtilis] Homo sapiens docking protein (DOK-2) mRNA, complete cds 1.7 2289097 [Cricetulus griseus] Llongillorum mRNA encoding calmodulin. >:: gb L18912 LILCALM ODU Lilium longiflorum selectivity mRNA (AF023270) probable			Rat mRNA for protein tyrosine	17.	1228035	is expressed ubiquitously.; The KIAA0191 protein retains the C2H2 zinc-finger at its N-	0.008
gene encoding troponin T isoforms. complete cds. Human peroxisome proliferator activated receptor mRNA, complete cds. 1.7 4033414 PUTATIVE IMPORTIN BETA-4 SUBUNIT Dendrobium crumenatum ACC synthase gene. complete cds 1.7 3122885 [Bacillus subtilis] Homo sapiens docking protein (DOK-2) mRNA, complete cds 1.7 2289097 [Cricetulus griseus] 184 AF034970 Lilongiflorum mRNA encoding calmodulin. > :: gb[L18912]LILCALM ODU Lilium longiflorum receptor mRNA (AF023270) probable	180	D30749	phosphatase		112000		
proliferator activated receptor mRNA, 182 L07592 complete cds. Dendrobium crumenatum ACC synthase gene. 183 U64031 complete cds Homo sapiens docking protein (DOK-2) mRNA, complete cds 184 AF034970 complete cds 185 Llongiflorum mRNA encoding calmodulin. > :: gb L18912 LILCALM ODU Lilium longiflorum mRNA encoding mRN	181	M15202	gene encoding troponin T isoforms.	1.7	731172		4e-04
Dendrobium crumenatum ACC synthase gene. 183 U64031 complete cds 1.7 3122885 [Bacillus subtilis] Homo sapiens docking protein (DOK-2) mRNA, complete cds 1.7 2289097 [Cricetulus griseus] 184 AF034970 complete cds 1.7 2289097 [Cricetulus griseus] 185 Liongiflorum mRNA encoding calmodulin. >:: gb L18912 LILCALM ODU Lilium longiflorum aclased line mRNA encoding mRNA encoding mRNA encoding mRNA encoding mRNA encoding mRNA encoding mRNA encoding mRNA encoding mRNA (AF023270) probable			proliferator activated receptor mRNA,	1.7	4033414		2e-06
183 U64031 complete cus 1.7			Dendrobium crumenatum ACC synthase gene.		3122885	SYNTHETASE synthetase	2e-11
L.longitlorum mRNA encoding calmodulin. > :: gb L18912 LILCALM ODU Lilium longiflorum calmodulin mPNA (AF023270) probable			Homo sapiens docking protein (DOK-2) mRNA, complete cds	1.7	2289097	alpha(1,3)fucosyltransferase	8e-12
	184	AP034970	L.longitlorum mRNA encoding calmodulin. > :: gb[L18912]LILCALM ODU Lilium longiflorum			(AF023270) probable	
185 Z12839 complete cds. 1.7 2511747 transcriptional regulator dre4	100	712830	1	1.7	2511747	transcriptional regulator dre4	4c-12

		The Contract of Co	anhank)	Negract Neight	oor (BlastX vs. Non-Redundant Pro	teins)
September 1	Nearest N	leighbor (BlastN vs. Go	enbank)	Mearest Neight	Joi (Blases VS. 1404)	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
					(270683) Weak similarity to	
		Equine arteritis virus (EAV) RNA genome > :: emb[A45589]A45589 Sequence 1 from Patent WO9519438 >	*		Human tyrosine-protein kinase CSK (SW:CSK_HUMAN); cDNA EST EMBL:C10908 comes from this gene; cDNA EST EMBL:C12822 comes from this gene; cDNA EST yk408c2.3 comes from this gene; cDNA EST yk408c2.5 Human tyrosine-protein kinase	
186	X53459	:: emb A58849 A58849 Sequence 1 from Patent WO9700963 > :: gb AR013959 AR013 959 Sequence 1 from patent US 5773235		3979817	CSK (SW:CSK_HUMAN); cDNA EST EMBL:C10908 comes from this gene; cDNA EST EMBL:C12822 comes from this gene; cDNA EST yk408c2.3 comes from this gene: cDNA EST yk408c2.5	le-14
		E. coli ddl gene encoding D-alanine: E alanine ligase and ftsQ and ftsA genes, complete cds, and		3870121	(Z70310) predicted using Genefinder; Similarity to Mouse ankyrin (PIR Acc. No. S37771); cDNA EST EMBL:T01923 comes from this gene; cDNA EST EMBL:D32335 comes from this gene; cDNA EST EMBL:D32723 comes from this gene; cDNA ES Genefinder; Similarity to Mouse ankyrin (PIR Acc. No. S37771); cDNA EST EMBL:T01923 comes from this gene; cDNA EST EMBL:D32335 comes from this gene; cDNA EST EMBL:D32723 comes from this gene; cDNA EST	
187	K02668	ftsZ gene, 5' end.	1.7	3879121	gene: cDNA ES	2e-19.
188	AB008375	Homo sapiens mRNA for osteoblast specific cysteine-rich protein. complete cds	e	2496945	HYPOTHETICAL 55.9 KD PROTEIN EEED8.6 IN CHROMOSOME II >gi 733603 (U23484) No definition line found [Caenorhabditis elegans]	Į.
189	L36603	Pseudomonas cepacii (clone Psudom70-1) heat shock protein 70 (hsp70) gene, complete cds	İ	2661842	(Y15732) DNA polymerase beta [Xenopus laevis]	a 6e-20

का र सर		The Way Co	-bank\	Nearest Neighl	oor (BlastX vs. Non-Redundant Pro	oteins)
÷×,,;;	Nearest N	leighbor (BlastN vs. Ge	noarik)	New Est Neight	1	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
	ACCESSION				HYPOTHETICAL 75.5 KD	
					PROTEIN C14A4.3 IN	T 1
					CHROMOSOME II	1 1
			İ		>gi 3874230 gnl PID e1351618	1
					protein (Swiss Prot accession	1 . 1
}		P.blakesleeanus		!	number P38376); cDNA EST	1 1
!	<u> </u>	mRNA GTP			yk220e10.5 comes from this	1
100	Z49760	cyclohydrolase I	1.7	1731181	gene [Caenorhabditis elegans]	3e-21
190	249700	Cyclonytholase 1			(AF125443) contains similarity	1 1
1		Human faity acid			to S. pombe phosphatidyl	1 1
	1	synthase gene, partial			synthase (GB:Z28295)	
191	U52428	cds	1.7	4226073	[Caenorhabditis elegans]	6e-25
191	032428	Human mitogen				1 1
1		induced nuclear				1
192	U12767	orphan receptor	1.6	<none></none>	<none></none>	<none></none>
172	012/0/	Orphia				1 1
i	l .	H. sapiens CpG DNA.	1		Į.	
1	1	clone 85a12, forward				
193	Z63478	read cpg85a12.ft1a.	1.6	<none></none>	<none></none>	<none></none>
122	2054.5	Homo sapiens				1.
1		inversin protein,	1	}		2707
194	AF084375	exons 8 and 9	1.6	<none></none>	<none></none>	<none></none>
1.	1200.000	Archaeoglobus				}
1	1	fulgidus section 165]			1
1	1	of 172 of the		ŀ		<none></none>
195	AE001114	complete genome	1.6	<none></none>	<none></none>	CHOILES
1	†	Homo sapiens		1		1
	ì	inversin protein.	j	_	NOVE	<none></none>
196	AF084375	exons 8 and 9	1.6	<none></none>	<none></none>	CHOINES
				1		
	1	Kluyveromyces lactis		I	l	l l
	ł	RNA polymerase II				
1		largest subunit gene,			<none></none>	<none></none>
197	U24217	partial cds	1.6	<none></none>	- CHOINES	13.0
	T	Helicobacter pylori	į.			
		26695 section 58 of			ĺ	1
		134 of the complete		NO.TO	<none></none>	<none></none>
198	AE000580	genome	1.6	<none></none>	- CHOILES	1 2,02

i fat is	Nearest N	leighbor (BlastN vs. Ge	nbank)	Nearest Neighb	or (BlastX vs. Non-Redundant Pro	teins)
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
	ACCESSION		<u> </u>			
					·	
		H.sapiens mRNA for	1		į	
		Drosophila female			i i	
		sterile homeotic				
		(FSH) homologue > ::				
		gb M80613 HUMFS			·	
	ł	HG Human homolog	,	İ		
	}	of Drosophila female				
		sterile homeotic mRNA, complete cds.	1.6	<none></none>	<none></none>	<none></none>
199	X62083	Plasmodium	1.0			
	i	brasilianum DNA				
	1	homologous to the	·	į		
	1	histidine-rich knob	<u> </u>			
	1	protein region of			O COCCUEN ODE V	
		Plasmodium	1		(M26647) ORF X [Saccharomyces cerevisiae]	8.4
200	M28064	falciparum.	1.6	457495	[Saccharolityces celeviside]	
		Streptomyces albus	1			Ì
	Ì	lipase precursor (lip)	ţ.	ł		ļ
	1	gene, complete cds, and unidentified 5'	1	1	(AC004877) sco-spondin-mucin-	1
	1	ORF and 3' ORF.	ļ		like; similar to P98167 uncertain	1
	*******	partial cds.	1.6	3638957	[Homo sapiens]	7.8
201	U03114	partial cus.				l
	l	Strix varia oocyte			DE DECENTOR	1
l	1	maturation factor			VITAMIN D3 RECEPTOR	1
		Mos (c-mos) proto-	l		(VDR) receptor [Rattus	6.4
202	U88422	oncogene, partial cds	1.6	137618	norvegicus]	
		Human pulmonary				}
	1	surfactant-associated			1	
	ł	protein SP-A			(Z38112) E03A3.6	
		(SFIP1) gene, complete cds.	1.6	3875423	[Caenorhabditis elegans]	4.9
203	M68519	Homo sapiens	 -			
1	1	transcription factor		1	GABA transport protein -	4.7
20	4 AF044575		1.6	2133625	tobacco hornworm	4.1
120	7004-375	Homo sapiens				1
1	1	(subclone 3_e10 from	n		(AJ005588) 5-epi-aristolochene	.]
1	1	P1 H21) DNA	ł	2/92207	synthase	4.6
20	5 L48476	sequence.	1.6	3687297	(Z81133) Similarity to Human	1
			ł		mRNA product KIAA0077	
1	1	Rat CNS 2',3'-cyclic		1	(TR:Q14997) [Caenorhabditis	
1		nucleotide 3-	1.6	3880315	elegans)	3.7
20	6 M18630	phosphodiesterase	1.0			

· · · · · · ·	Nearest N	leighbor (BlastN vs. Go	nbank)	Nearest Neight	oor (BlastX vs. Non-Redundant Pro	iteins)
SEQ	14carest 1	icignoor (3.				DALATIE
Ð	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	PVALUE
_	11000001011					
		Arabidopsis thaliana				
		cellulose synthase				
		catalytic subunit (Ath-				
		B) mRNA, complete			TUMOR-ASSOCIATED	3.6
207	AF027174	cds	1.6	267068	ANTIGEN L6	3.0
	714 02 1 1 1	Babesia microti heat			(U53155) strong similarity to the carboxyl two-thirds of valyl-	}
		shock protein 70		•	the carboxyl two-thirds of varyl-	
		(hsp70) gene,			tRNA synthetases [Caenorhabditis elegans]	2.2
208	U53448	complete cds	1.6	1255429	PROBABLE	
			}		SERINE/THREONINE-	ļ
			1		PROTEIN KINASE CY49.28	1
	1	Homo sapiens] .]		>gi 1370255 gnl PID e247094	l
		inversin protein		1720076	(Z73966) pknJ	1.2
209	AF084367	mRNA, complete cds	1.6	1730076	(2/3900/ pkil)	
		Yeast dis 1+ gene for			(AF010496) maltose transport	!
	1	p93dis1, complete	1	3128353	inner membrane protein	1.2
210	D55635	cds 2	1.6 .	3128333		
	1	Streptomyces sp. 2-		j	(X83413) DR5 [Human	
]	dehydro-3-		j	herpesvirus 6] >gi 853972	ļ
		deoxyphosphohepton	ļ		(X83413) DR5 [Human	
	. 7025756	ate aldolase gene.	1.6	853971	herpesvirus 6]	0.97
211	AF035756	Ocuniculus rPTPA	1.5		(Y17034) Bassoon [Mus	
212	X73479	mRNA	1.6	3413810	musculus]	0.94
212	A/34/9	HIRCHA			(U95142) putative G-protein-	1
	1		į.		coupled receptor G-protein-	
		H. sapiens mRNA for			coupled receptor [Arabidopsis	0.73
213	X98330	ryanodine receptor 2	1.6	2072986	thaliana]	
	1,,,,,,,,,	7			NECDIN >gi 91129 pir JN0148	
	ì		i	ţ	necdin, brain - mouse	1
		Panserina FMR1	1		>gi 200020 (M80840) necdin	0.42
214	X64194	gene exons 1 and 2	1.6	128014	[Mus musculus]	1
	1	Caenorhabditis		T.		ľ
	1	elegans cosmid	1		1	
	I	F53B8, complete	}	1	(U23517) D1022.7	
		sequence	1		[Caenorhabditis elegans]	1
	1	[Caenorhabditis	1	746516	>gi 3258651 elegans	0.19
215	Z92788	elegans]	1.6	740310		7
	ı	Methanobacterium			INTERFERON-ALPHA/BETA	4
l	ł	thermoautotrophicum	.		RECEPTOR ALPHA CHAIN	
	1	from bases 1098908			PRECURSOR (IFN-ALPHA-	
		to 1112186 (section	1	1	REC) >gi 346520 pir S27387	1
	1	94 of 148) of the			interferon alpha receptor type 1	
			1.6	462415	bovine >gi 432	0.001
21	6 AE000888	Leombiere Senome				

E.G.	Nearest N	eighbor (BlastN vs. Ger	nbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant Pro	terns,
EQ		DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
<u> </u>	ACCESSION	DESCRIPTION OF THE PROPERTY OF				
					HYPOTHETICAL 55.9 KD	
ı		Homo sapiens mRNA	ŀ		PROTEIN EEED8.6 IN	
1		for osteoblast specific	l		CHROMOSOME II >gi 733603	
- 1		cysteine-rich protein.	İ		(U23484) No definition line	
[. 5000055	cysteine-rich protein.	1.6	2496945	found [Caenorhabditis elegans]	le-18
217	AB008375	complete cds			(Z70750) similar to vanadate	
- 1			i	i	resistance protein	}
ı		Orang-utan involucrin			transmembranous domains	2.26
	3525212	gene, complete cds.	1.6	3875131	[Caenorhabditis elegans]	3e-26
218	M25312	Cyprinus carpio				1
		mRNA for MyoD,		,	NO.	<none:< td=""></none:<>
219	AB012882	complete cds	1.5	<none></none>	<none></none>	OTOLIE.
217	AD012002	Caenorhabditis	•			1
		elegans cosmid			<none></none>	<none:< td=""></none:<>
220	U29487	C09C7	1.5	<none></none>	SHOWES	1
				1	integral membrane protein -	ŀ
	1		Ì		Streptomyces pristinaespiralis	1
	ł		1		>gi 872306 (X84072) integral	
	l			Į.	membrane protein	
	į	M.musculus mRNA	١, , ,	1364094	(Streptomyces pristinaespiralis)	4.3
221	X74760	for Notch 3	1.5	1304094	TEXOGLUCANASE II	
			Ì		PRECURSOR cellulose 1,4-bet	al
	l .				cellobiosidase (EC 3.2.1.91) II	
	1		į.	1	precursor - fungus (Trichoderm	a
	1	Lycopersicon esculentum class II	1		reesei) 1,4-beta-cellobiosidase	1
	1	small heat shock		1	(EC 3.2.1.91) 11 - fungus	}
1	l l	protein Le-HSP17.6	}		cellobiohydrolase II	1
	,,,,,,,,,	mRNA, complete cds	1.5	121855	[Trichoderma reesei]	4.3
222	U72396	micha, complete cus	+			
	I	Human myosin-IXb	1			4.2
223	U42391	mRNA, complete cds	1.5	3688428	(AJ011534) sucrose synthase	
-223	042391					
1		Pongo pygmaeus	1	1		
	1	gamma-1 and gamma	1-	1	(M13144) inhibin A [Homo	
	1	2 globin genes.		125.12		0.22
224	M92296	complete cds.	1.5	186413	sapiens] (AF029791) UDP-	
			1	1	Gal:betaGlcNAc beta 1,3-	-
1	1				galactosyltranferase-II [Mus	
1	1	C.japonica mRNA fo	OT .	2745737	musculus	3e-08
225	5 X94144	QNR-71 protein	1.5	2143131	11143-61-603	

				Manust Neighl	oor (BlastX vs. Non-Redundant Pro	teins)
100	Nearest N	eighbor (BlastN vs. Ge	nbank)	Nearest Neigh	T T	
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
	ACCESSION					
					(X90568) Protein sequence and	
ŀ					annotation available soon via	
1			i		Swiss-Prot; available at present	
1		Homo sapiens mRNA	l		via e-mail from	
1		for KIAA0657			LABEIT@EMBL-	4e-13
۰.,		protein, partial cds	1.5	1212992	Heidelberg.DE [Homo sapiens]	46-13
226	AB014557	Borrelia burgdorferi				
1		oligopeptide				1
	ĺ	permease homolog				
		OppAIV (oppAIV)				<none></none>
207	AF000948	gene, complete cds	1.3	<none></none>	<none></none>	CHOINES
227	Ar000948	Mus musculus			MYC PROTO-ONCOGENE	
		RAB/Rip protein			PROTEIN (C-MYC) proto-	2.6
228	AF057287	mRNA, partial cds.	1.3	2498005	oncogene [Sus scrofa]	2.0
220	AF031281	Drosophila				}
		melanogaster	İ			
	i	vacuolar ATPase	1		NONE	<none:< td=""></none:<>
229	U38951	subunit E	1.1	<none></none>	<none> (U90209) RNA polymerase II</none>	1210112
	1 030,31	Homo sapiens			largest subunit [Bonnemaisonia	1
		myogenic	į.			2.3
230	AF027148	determining factor 3	1.1	3172134	hamifera]	
230	1	Mus musculus histon	е	1		}
	i	deacetylase 3	}		1	1
		(Hdac3) gene, exons	1	1	(U66220) unknown	Ì
		4 through 15 and			[Nannocystis exedens]	0.25
231	AF079310	complete cds	1.0	1657601	(X91638) BRM protein [Gallus	
<u> </u>		P.radiata lac gene for		996020	gallus]	0.31
232	X52134	laccase	0.95	996020	ganus	7
		Human mRNA for	l l			1
[1	Neuroblastoma,	0.03	<none></none>	<none></none>	<none< td=""></none<>
233	D89016	complete cds	0.93	CNONES	(AL022238) dJ 1042K10.2.1	
<u> </u>		C.familiaris VIP36	1 .	,	(novel protein with probable	1
	[·	(vesicular integral-	.		rabGAP domains and Src	
	1	membrane protein of	1 000	4176446	homology domain 3)	7e-81
234	4 X76392	36 kDa) mRNA	0.93	41/0440		
		Mus musculus	ł			}
1	ł	Pontin52 mRNA.	0.90	<none></none>	<none></none>	<none< td=""></none<>
23.	5 AF100694	complete cds	1. 0.90	41.01.10		***

				No. No. abb	oor (BlastX vs. Non-Redundant Pro	teins)
444	Nearest N	leighbor (BlastN vs. Ge	nbank)	Nearest Neight	OF (Blasta Vs. Non-Reddildant 1)	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
<u> </u>	ACCESSION				EGIZ PROTEIN PRECURSOR	
					(EARLY GI TRANSCRIPT 2)	
1			i		>gi 1362345 pir S55862	
1					probable membrane protein	1
ļ			1		YNL327w - yeast	i i
}					(Saccharomyces cerevisiae)	1
l		Archaeoglobus	l		cerevisiae]	1
1		fulgidus section 116	1		>gi 1302445 gnl PID e239572	{
1	1	of 172 of the	1		(Z71603) ORF YNL327w	
236	AE000991	complete genome	0.90	1176579	[Saccharomyces cerevisiae]	6.9
230	AL000771	S.cerevisiae]
1		chromosome II	,		·	1 1
j]	reading frame ORF				NONES
237	Z35922	YBR053c	0.86	<none></none>	<none></none>	<none></none>
1-25	-					1 1
1	1	Rattus norvegicus			1	1
1	l .	metabotropic	İ		(manages) to seek seized septein	
1	1	glutamate receptor 4b	i i		(Z80225) hypothetical protein	4.1
238	U47331	mRNA, complete cds.	0.82	1550703	Rv2662	 4.1
1					ļ	1
1	1	H.sapiens 1g germline	}		(AF052587) F14 [Xylella	1
1		kappa-chain gene			1	6.7
239	X72810	variable region (L3)	0.69	3023063	fastidiosa]	
	1		1			
	1	Escherichia coli	į.	ţ		
1	1	genes faeG, faeH,	Į	1		1
1	Ĭ	fael, fael and IS629-				
1	i i	like insertion	į	1		1
l	ì	sequence. > ::	i	1		
	1	emb Z11710 ECFAE	1		į.	1
ļ		HU E.coli faeH, faeI	<u> </u>			1
1.	1	and fael genes	1	· ·	(AC002338) laccase isolog	1
	1	encoding FaeH, Fael	0.00	2347188	[Arabidopsis thaliana] thaliana]	3.9
240	Z11700	and FaeJ proteins	0.69	254/100	[/ Hadidopsis Historia, Historia	1
	1	Phrynosoma	-	ļ	1	1
	1	douglassii NADH	}	1		
	1	dehydrogenase	1	}		1
	1	subunit 4 (ND4)	Į.	1	1	
1	ł	gene, mitochondrial	·	1		
1.	1	gene encoding	1			
	I	mitochondrial	0.65	<none></none>	<none></none>	<none></none>
24	U71597	protein, partial cds	0.65	1 010112		

			-ble)	Nearest Neighl	oor (BlastX vs. Non-Redundant Pro	teins)
	Nearest N	eighbor (BlastN vs. Ge	ndank)	rearest riorgan		
EQ		DESCRIPTION	P VALUE	ACCESSION_	DESCRIPTION	P VALUE
	ACCESSION	DESCRIPTION			GLUTAMYL-TRNA	
					SYNTHETASE glutamate	
- 1			ł		tRNA ligase (EC 6.1.1.17) -	
					Haemophilus influenzae (strain	
			i		Rd KW20) >gi 1573240	
		Ammonia species			(U32713) glutamyl-tRNA	
		LSU rRNA gene			synthetase (gltX) [Haemophilus	
	l	(partial; isolate Tr S	0.64	1174506	influenzae Rd]	1.2
242	Z77798	5: clone 16)	0.64	11/4300	Timec.izae esa	
		Day A Co.		ı		1
]	Human mRNA for			ultra-high-sulfur keratin 1 -	
	1	golgi antigen gcp372,	0.64	111230	mouse	1e-05
243	D25542	complete cds	0.64	111230		
	l				(Z99709) similar to Elongation	
	ł			ļ.	factor Tu family (contains	1
	· I	1	}		ATP/GTP binding P-loop);	Ì
	1 .	ł	}	ł	cDNA EST EMBL: D76223]
	1			Ī	comes from this gene; cDNA	
		Cow dopamine		1	EST vk478c5.5 comes from this	
	1	transporter mRNA,	0.64	3874972	gene [Caenorhabditis elegans]	8c-06
244	M80234	putative cds.	0.64	3674772	EPIDERMAL GROWTH	
			1	ł	FACTOR RECEPTOR	
	1	ļ	Į.	§ .	KINASE SUBSTRATE EPS8	j
			}	1	>gil530823 (U12535) epiderma	1
ı		Homo sapiens mRNA	`[i	growth factor receptor kinase	1
		for KIAA0449	0.64	2833239	substrate [Homo sapiens]	2e-14
245	AB007918	protein, partial cds	0.64	20,52,55		
		Human U266	1	i		1
1	İ	rearranged DNA for	1		(U95102) mitotic	
1	1	lambda-	1	1	phosphoprotein 90 [Xenopus	ļ
	Į.	immunoglobulin ligh		2072301	laevisl	1.5
246	X51754	chain	0.63	2072301	IMC VIST	
		Helicobacter pylori,		1	-	
1	ı	strain 199 section 115	'l		1	1
1.	1.	of 132 of the	0.00	<none></none>	<none></none>	<none< td=""></none<>
247	7 AE001554	complete genome	0.62	CHOINES		
	1	H.sapiens CpG DNA		1	1	
	1	clone 96e7, reverse	"[1 _		
_			0.62	<none></none>	<none></none>	<none< td=""></none<>
24	8 Z64067	read cpg96e7.rtla. Pinus sylvestris	+	1		
1		microsatellite DNA,	ì			
		Imicrosatenne Diva-	1	<none></none>	<none></none>	<none< td=""></none<>

	(BlastX vs. Non-Redundant Pro	Nearest Neighb	ank)	eighbor (BlastN vs. Genl	Negreet N.	i e
VALUE	DESCRIPTION	ACCESSION	VALUE			EQ
			VALUE	DESCRIPTION I	CCESSION	D A
	PHOTOSYSTEM II 10 KD					
	PHOSPHOPROTEIN deltoides]		1			
	>gi 2143326 gnl PID e319090		1	1	1	ı
	(Y13328) 10kDa		l		1	- 1
	phosphoprotein [Populus		į.			1
7.9	deltoides	2493689	0.00	Bacteriophage P1 ban		- 1
		2473003	0.62	gene	AJ011592	250
				Xenopus laevis		
	(AL034463) putative single-			survival of motor		1
	strand polynucleotide binding			neuron protein		- 1
	protein [Schizosaccharomyces			interacting protein 1		1
2.0	nombel	4007790	0.72	(SIP1) mRNA.		
	(U40763) CARS-Cyp [Homo	4007770	0.62	complete cds	AF027151	251
0.90	ilensl	1117968	0.63	Helobdella triserialis		
	(AC005058) similar to calcium-	111/700	0.62	mRNA for actin	AJ000376	252
}	independent phospholipase A2;					
[similar to AC004392			\ . \		- 1
	(PID:g3367519) [Homo					- 1
6e-51	sapiens]	4176370	0.62	Rat thymosin beta 4		
		11.05.0	0.02	gene (pTB4G).intron.	M69231	253
l	}			Homo sapiens X11L2 mRNA for X11-like		
<none:< td=""><td></td><td></td><td></td><td>mRNA for All-like</td><td></td><td></td></none:<>				mRNA for All-like		
SNONE	<none></none>	<none></none>	0.61	protein 2, complete		
	· ·			cds Bacteroides	AB021638	254
				gingivalis DNA for	l	
ì				arginyl	l	
<none:< td=""><td></td><td></td><td></td><td>endopeptidase.</td><td>]</td><td></td></none:<>				endopeptidase.]	
CATOLAL	<none></none>	<none></none>	0.61	complete cds	D06470	
<none< td=""><td>NONES</td><td></td><td></td><td>A.thaliana ATPase</td><td>D26470</td><td>255</td></none<>	NONES			A.thaliana ATPase	D26470	255
1 210	<none></none>	<none></none>	0.61	gene, complete cds.	J04737	
1	1			Bos taurus clone	304/3/	256
	(Y09905) snail like protein	}	1	bm1308	1	i i
0.51	(Y09905) shall like protein	'		microsatellite and are	1	
+	[Gallus gailus]	1922280	0.61	lp repeat region.	U06756	257
1		1			000730	257
			1	p15=cyclin D-		1
	}	1	1	dependent kinases 4	1	1
}	1	1	.	and 6-binding	1	1
1	hypothetical protein 253 -	1		protein/p15 product	•	1
1	Streptomyces griseus plasmid	I		{exon/intron 1}	1	1
0.13	Suchiomaces Brisens brasing		,	(human, brain tumors	1	1
	pSG1 (fragment)	484938	0.61	1	\$75756	250
1	(Z70750) similar to vanadate	Į.		Drosophila	, 3,3,30	258
1	resistance protein		1	melanogaster tumor	1	
	transmembranous domains			supressor (warts)	1	1
1e-09	[Caenorhabditis elegans]		1	mRNA exons 1-8.	1	1
	[Caenornavorus ciceans]	3875131	0.61	1	D L39837	259

1.01 W	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
非种类	Nearest N	Neighbor (BlastN vs. G	enbank)	Nearest Neight	BOI (BIASIX VS. INON-REGULARIAN I II	Jienis)	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					(AF125443) contains similarity		
		Human fatty acid			to S. pombe phosphatidyl		
		synthase gene, partial	1		synthase (GB:Z28295)	1 · 1	
260	U52428	cds	0.61	4226073	[Caenorhabditis elegans]	2e-26	
		Plasmodium					
)		falciparum gene for			1	1	
1 1		heat-shock protein				1	
261	X15292	pPf203	0.60	<none></none>	<none></none>	<none></none>	
1 1		Homo sapiens mRNA				· •	
		for KIAA0856			(U00043) No definition line		
262	AB020663	protein, partial cds	0.60	470341	found [Caenorhabditis elegans] GALACTOSE-BINDING	5.7	
					PROTEIN REGULATOR		
<u> </u>					glucose/galactose binding	1	
					protein regulator -	1	
,					Agrobacterium tumefaciens	1	
ļ					>gi 142228 (L10424)	1	
		Human checkpoint			glucose/galactose binding		
		suppressor I mRNA.	0.40	644376	protein regulator	5.7	
263	U68723	complete cds	0.60	544375	protein regulator	 	
1		S.griseus sporulation					
		protein genes 1590			(AF012871) Mergla' [Mus		
344		į.	0.60	2582017	musculus	3.3	
264	M32687	and 1422. Homo sapiens	0.00	2502017	mascarati		
		NKCC2 gene, exon 4,			(AF010496) maltose transport	1	
265	AJ005331	isoform B	0.60	3128353	inner membrane protein	1.5	
203	ונפנטטנע _	Mus musculus RGL					
		protein mRNA,			(U90533) serine protease		
266	U14103	complete cds.	0.60	4099845	inhibitor [Streptomyces fradiae]	0.098	
1-50		Xenopus laevis XL-					
		INCENP (XL-		•	1	1	
		INCENP) mRNA,			(AF047897) ankyrin-like protein		
267	U95094	complete cds	0.59	3282851	HGE-ANK [Ehrlichia sp. BDS]	5.5	
	T	Methanobacterium					
	Į	thermoautotrophicum					
1	S	from bases 896604 to					
	i l	912784 (section 78 of			HYPOTHETICAL 24.5 KD		
	1	148) of the complete			PROTEIN IN NADB-SRMB		
268	AE000872	genome	0.59	401553	INTERGENIC REGION	4.3	

70.3	Nearest N	Veighbor (BlastN vs. Ge	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					nypomencai protein - human		
		· · · · · · · · · · · · · · · · · · ·			herpesvirus 4 reading frame 1		
					[Human herpesvirus 4] 2		
! !					[Human herpesvirus 4]	!	
1 1					>gi 1334838 gnl PID e25079 4		
l '					[Human herpesvirus 4]		
					>gi 1334840 gn1 PID e25081 6		
					[Human herpesvirus 4]		
1					>gi 1334842 gn1 PID e25067 8		
1					[Human herpesvirus 4]	}	
		Gallus gallus achaete-	5-		>gi 1334844 gnl PID e25069 10	į	
}		scute homologue		•	[Human.herpesvirus 4]	1	
}		(ASH) mRNA,			>gi 1334846 gnl PID e25071 12		
269	L11871	complete cds.	0.59	628110	[Human herpesvirus 4]	4.2	
					NITROGENASE IRON-IRON		
1					PROTEIN ALPHA CHAIN		
					INITROGENASE		
		Oryctolagus			COMPONENT D	1	
i .		cuniculus glycogen			(DINITROGENASE) capsulatus		
1	i	synthase mRNA,			>gi 312238 (X70033)		
270	AF017114	complete cds	0.59	728856	alternative nitrogenase	2.4	
1	1201/121						
	,	Homo sapiens beta-			(AF067155) truncated rev.		
	}	casein (CSN2) gene,			protein [Human	٠, ا	
271	AF027807	complete cds	0.59	3252932	immunodeficiency virus type 1]	1.5	
		Human Wnt10B			(Z67990) similar to cuticle		
272	U81787	mRNA, complete cds	0.59	3875538	collagen	1.4	
212	081/8/	Apteryx australis 16S					
	1	ribosomal RNA gene.					
	1	mitochondrial gene					
1]	for mitochondrial	}		(AF055088) ATP-binding		
].		RNA, partial		•	cassette, PsaB [Streptococcus		
273	U76036	sequence	0.59	4193356	pneumoniae]	0.83	
					PTB-ASSOCIATED SPLICING		
					FACTOR (PSF) long form -		
1		Homo sapiens mRNA			human >gi 38458 (X70944)		
		for KIAA0664			PTB-associated splicing factor		
274	AB014564	protein, partial cds	0.59	1709851	[Homo sapiens]	0.17	
		Homo sapiens cyclin-					
1	1	dependent kinase			1		
1		inhibitor 2D			(AL032626) Y37D8A.17	}	
		(CDKN2D) gene.	0.50	2025212	T	3e-10	
275	AF044171	partial cds	0.59	3925213	[Caenorhabditis elegans]	30-10	

				Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
-12	Nearest N	leighbor (BlastN vs. Ge	nbank)	Nearest Neighbor (Blasta Vs. 1701-Reduction 1.151-15)				
SEO				·				
D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
	ACCEDOIO							
		Saccharomyces				1		
i i		cerevisiae cdc2/cdc28				1		
		related protein kinase			(Z81130) T23G11.9			
276	L19640	gene, complete cds.	0.59	3880115	[Caenorhabditis elegans]	1e-21		
2/0	£19040	Human DNA				i l		
1		sequence from			1	i d		
		cosmid E140G5 on			1	l 1		
,		chromosome 22,				<u> </u>		
Ì	,	complete sequence				NONTE		
277	Z80999	[Homo sapiens]	0.58	<none></none>	<none></none>	<none></none>		
		H.sapiens WNT8B			1,015	<none></none>		
278	Y11108	gene	0.58	<none></none>	<none></none>	ANONE		
	l	Sphyraena idiastes	·		1	1 1		
1	1	lactate dehydrogenase			NONTE	<none></none>		
279	U80001	Α	0.58	<none></none>	<none></none>	CHOILES		
		S.cerevisiae	}	,		1		
1	<u> </u>	chromosome X		l		1 1		
1	ł	reading frame ORF	1		<none></none>	<none></none>		
280	Z49637	YJR137c	0.58	<none></none>	A TONE	2.0		
		H.sapiens ALAD						
1		gene for	1			1		
	İ	porphobilinogen	2	NONE	<none></none>	<none></none>		
281	X64467	synthase	0.58	<none></none>	1 1101.25			
	1	G.gallus hox B3	0.58	<none></none>	<none></none>	<none></none>		
282	X74506	mRNA	0.38	(110/12)				
1		Cochliobolus	1	1	l .	1		
1		heterostrophus	0.58	<none></none>	<none></none>	<none></none>		
283	U68040	polyketide synthase	0.38	4.0.1.5		1		
1			1		1			
1	1	Arabidopsis thaliana				[
ł		putative auxin efflux	1	Ì	1	ł		
1	1	carrier protein (PIN1	,	1 .		i _		
204	A ECOCOR 4			<none></none>	<none></none>	<none></none>		
284	AF089084	Rattus norvegicus	1 222	1		1		
		ROK-alpha mRNA,	1		1			
285	U38481	complete cds	0.58	<none></none>	<none></none>	<none></none>		
20-	030-01					1		
	,	Homo sapiens G	1	1				
1	ļ	protein beta 5 subuni	t	1	(AC004684) hypothetical	9.2		
286	AF017656			3236249	protein [Arabidopsis thaliana]	7.2		
1-20	1	Human glutathione			(U55366) Similar to cuticle			
	1	transferase class mu			collagen [Caenorhabditis	7.1		
28	M96234	number 4	0.58	1280073	elegans]	 		
		Human mRNA for	1		(U28741) F35D2.1 gene	1		
		KIAA0341 gene.	1	041222	product [Caenorhabditis	7.1		
28	8 AB002339	partial cds	0.58	861293	elegans]			

- Article	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	iveatest i	tergitoor (Diasar vs. C	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Neisseria gonorrhoeae carbamoyl phosphate synthetase (glutamine) small			(AF020283) DG2044 gene	·	
		subunit (carA) and large subunit (carB)	0.50	2425135	product [Dictyostelium discoideum]	5.3	
289	U11295	genes, complete cds. Human mRNA for	0.58	2423133	(U49836) gamma-glutamyl	 	
290	D80001	KIAA0179 gene.	0.58	4097223	transpeptidase precursor [Brugia malayi]	4.1	
-		Escherichia coli genes faeG, faeH, faeI, faeJ and IS629- like insertion sequence. > :: emb Z11710 ECFAE HIJ E.coli faeH, faeI					
291	Z11700	and faeJ genes encoding FaeH, FaeI and FaeJ proteins	0.58	2347188	(AC002338) laccase isolog [Arabidopsis thaliana] thaliana]	3.2	
292	M77350	Mouse hair keratin A1 (MHKA1) gene, complete cds.	0.58	141165	HYPOTHETICAL 8.3 KD PROTEIN >gi 62179 (AB004461) DNA polymerase	3.2	
293	X63787	T.thermophila gene for snRNA U3-2	0.58	2826900	alpha catalytic subunit [Oryza sativa]	3.1	
294	D63881	Human mRNA for KIAA0160 gene, partial cds	0.58	1934730	(U95036) germin-like protein [Arabidopsis thaliana]	3.1	
295	U39378	Gymnocarena mexicana 16S ribosomal RNA gene mitochondrial gene encoding mitochondrial RNA, partial sequence	0.58	2194131	(AC002062) Similar to Synechocystis antiviral protein	3.1	
296		P.pastoris PRC1 gene > :: dbj E12103 E12103 DNA encoding precursor of protease from Pichia pastoris		3914197	OCCLUDIN >gi 1276983 (U49221) occludin [Canis familiaris] >gi 1589181 prf 2210347D occludin [Canis familiaris]	3.1	

				Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
- 	Nearest N	eighbor (BlastN vs. Ge	nbank)	Nearest Neight)		
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
<u> </u>	ACCESSION						
		A.thaliana (L.Heynh.)				l l	
		chloroplast mRNA			(D38529) DRPLA protein	1	
		for recombinant APS-	II.		(D38529) DRPLA protein	2.4	
297	X75782	kinase	0.58	1732444	[Homo sapiens]		
291	A 3702	Mouse platelet-		1		1	
		derived growth factor		I	·	1	
]	B chain musculus		İ	1		
	1	platelet-derived				1	
	ł .	growth factor beta-			(AF055985) pyrrolidone-rich		
	ł .	chain (sis) gene, exon		. 3025832	antigen (Onchocerca volvulus)	1.4	
298	M64848	5.	0.58	. 3023832			
		Helicobacter pylori.		1	.		
		strain J99 section 21		1	(AF037454) ubiquitin protein	1	
	1	of 132 of the	0.58	2827198	ligase [Mus musculus]	1.1	
299	AE001460	complete genome	0.58		CHOI PROTEIN	1	
			Ĭ.		>gi 320737 pir S30818		
	1		1		hypothetical protein YER164w	1	
	I	1		1	yeast (Saccharomyces	}	
	1	M.musculus gene for	Ĭ	1	cerevisiae) >gi 603404		
}	į	protein kinase C-	1	1	(U18917) Chd1p: transcriptions	*1	
1	1	gamma (exon1 and	1	1	regulator [Saccharomyces	1.1	
200	X65720	exon 2)	0.58_	418395	cerevisiae] SEX-DETERMINING	+	
300	A03720	CRON 27			REGION Y PROTEIN		
1	1	Arabidopsis thaliana	1		determining protein [Mus	0.62	
30	1 AF043130	lactate dehydrogenas	e 0.58	3024637	determining protein (1.245		
130	1 12 0 13 13 1	Human genes for	•	į .		1	
1		collagen type IV	1		(U64835) T09D3.3	İ	
	-1	alpha 5 and 6, exon	1	1458250	[Caenorhabditis elegans]	0.36	
30	2 D28116	and exon 1'	0.58	1438230			
		Archaeoglobus	اء	1	1	1	
		fulgidus section 32 d	21	•	(Z97991) hypothetical protein		
1.	1.	172 of the complete	0.58	2276333	Rv0336	0.36	
30	3 AE00107	5 genome				1	
	1			1		1	
1	1	Rhodococcus opacu	s	1		1	
1	l l	chloromuconate	1	1		1	
	1	cycloisomerase	ļ	1	mucin 7 precursor, salivary -	1	
1		transposase homolo	g			0.28	
١,,	04 AF00394	18 genes, complete cd	0.58	477072	human		
٣	11 0037	Human MAGE-7]			
1	1	antigen (MAGE7)			HOMEOBOX PROTEIN HO	x-	
	1	pseudogene, compl	ete	2202050	CII	0.054	
1.	05 · U1069	2 cds	0.58	3287858	1011		

				Nearest Neight	oor (BlastX vs. Non-Redundant Pro	teins)
447	Nearest N	leighbor (BlastN vs. Ge	noank)	Mentest treight	,	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
	ACCESSION					
1		Rhodococcus opacus				
	ł	chloromuconate	1		1	
1		cycloisomerase				
	1	transposase homolog			(AF058803) mucin 4 [Homo	0.041
-06	45003040	genes, complete cds	0.58	3551821	sapiens]	0.041
306	AF003948	H.sapiens HFH4			VAV PROTO-ONCOGENE	
	1	gene, exon 1 and			>gi 55221 (X64361) proto-	0.024
	V00250	joined CDS	0.58	137483	oncogene [Mus musculus]	0.024
307	X99350	Homo sapiens mRNA				i .
		for Ig heavy chain	1		DOGGAS Offices	1
	I .	variable region, clone	l	z.	(AC003682) R27945_2 [Homo	0.018
200	AJ234282	lc	0.58	3264846	sapiens]	0.018
308	MJ234202	Mus musculus histone		1	1	
,	1	deacetylase 3	1			į
İ	1	(Hdac3) gene, exons	i			
1	t	4 through 15 and			(U66220) unknown	0.014
309	AF079310	complete cds	0.58	1657601	[Nannocystis exedens]	1-0.514
303	7407350	Human thiopurine		ł	(AF063020) lens epithelium-	1
	1	methyltransferase	į		derived growth factor [Homo	
l	1	(TPMT) gene, exons				0.011
310	AF019367	6 and 7	0.58	3283352	sapiens]	
 		M.musculus gene for			(U38291) microtubule-	1
l		protein kinase C-	i		associated protein la [Homo	}
1	ł	gamma (exon1 and		.700078	sapiens)	0.008
31	X65720	exon 2) .	0.58	1790878	Sapicitaj	
			. i	1	1	1
1	1	Homo sapiens mRN	*	ì	SYNAPSINS IA AND IB	1
1		for KIAA0583	0.50	1351166	>gi 163713	0.006
31	2 AB011155	protein, partial cds	0.58	1331100		
				1	(D84307) phosphoethanolamin	e
1	1			. .	cytidylyltransferase [Homo	ł
1	1	H.sapiens mRNA for	0.58	1817548	sapiens]	0.001
31	3 X63692	DNA	0.00	101,545		
	i	Feline	i	1		
		immunodeficiency	-	ļ]
	1	virus isolate FIV-	1	1		1
	l l	Pco336-8 pol		1	(U93872) ORF 73, contains	
1	l .	polyprotein (pol)	0.58	2246532	large complex repeat CR 73	2e-05
31	4 U53746	gene, partial cds	- 0.58-			1
1	1	Ratius norvegicus	_		(M64793) salivary proline-rich	1
-	i i	(clone rt1-1) pseudo	0.58	206712	protein [Rattus norvegicus]	1e-05
3	15 K00436	Gly-tRNA gene.				

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
i							
		HSF2=heat shock					
	,	factor 2 {alternatively					
-		spliced, splice	: I		1		
	•	junction region}					
		[mice, CBA/J, testis,	}		(A 10000000) -DET1		
		Genomic, 120 nt.	0.50	4038594	(AJ222798) tDET1 protein [Lycopersicon esculentum]	3e-06	
316	S79632	segment 2 of 3]	0.58	4038394	(U55376) coded for by C.	36-00	
					elegans cDNA cm21e6; coded		
l					for by C. elegans cDNA		
1 1					cm01e2; similar to melibiose		
}				•	carrier protein		
] [Rat liver mRNA for			(thiomethylgalactoside permease		
317	D43964	Kan-1, complete cds	0.58	1280135	II)	1c-08	
			_		EPIDERMAL GROWTH		
!!			•		FACTOR RECEPTOR KINASE SUBSTRATE EPS8		
1 1					>gi 530823 (U12535) epidermal		
		Homo sapiens mRNA for KIAA0449			growth factor receptor kinase		
318	AB007918	protein, partial cds	0.58	2833239	substrate [Homo sapiens]	3e-13	
318	AB00/918	Homo sapiens mRNA	- 0.50	2033237	0003000 (210110 0000)		
1		for Efs1, complete			(D45027) 25 kDa trypsin		
319	AB001466	cds	0.58	2943716	inhibitor [Homo sapiens]	2e-14	
		Saccharomyces					
1 1	\	cerevisiae IRE1 gene	1				
		for putative protein			(Z81130) T23G11.9	021	
320	Z11701	kinase.	0.58	3880115	[Caenorhabditis elegans] (Z83819) dJ146H21.2 (similar	9e-21	
1 1		S.cerevisiae chromosome X			to CYTOCHROME B-245		
1		reading frame ORF			HEAVY CHAIN) [Homo	1	
321	Z49535	YJR035w	0.58	4106562	sapiens]	3e-33	
221	£-7233	S.cerevisiae DBF20					
-322	M62506	gene, complete cds.	0.57	<none></none>	<none></none>	<none></none>	
		Yeast PSS gene for					
		phosphatidylserine					
323	X05944	synthetase	0.57	<none></none>	<none></none>	<none></none>	
		Snail gene for ADP-	1				
324	D20524	ribosyl cyclase,	0.57	<none></none>	<none></none>	<none></none>	
324	D38536_	complete cds S.cerevisiae	0.37	KNONES	110112	31.01.102	
		chromosome XV	. 1				
		reading frame ORF	I		1		
325	Z75004	YOR096w	0.57	<none></none>	<none></none>	<none></none>	
		Homo sapiens					
		(subclone 10_e10					
		from PI H16) DNA				ANONE:	
326	L77034	sequence.	0.57	<none></none>	<none></none>	<none></none>	

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		on all on C	-book) I	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	Nearest N	leighbor (BlastN vs. G	nbank)	Nearest Neighbor (2)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Cyprinus carpio c-					
		myc gene for c-Myc.			NONE	<none></none>	
327	D37887	complete cds	0.57	<none></none>	<none></none>	ANONES	
		Homo sapiens mRNA for KIAA0662			(M57576) Ig kappa chain [Mus	8.9	
328	AB014562	protein, partial cds	0.57	197406	musculus]		
		Human DNA sequence from cosmid L75B9, Huntington's Disease Region, chromosome		•	chaperonin containing TCP-1 complex gamma chain - African clawed frog >gi 793886	8.9	
329	Z69651	4p16.3	0.57	1079280	(X84990) Cctg	0.5	
220	D00005	Mesocricetus auratus mRNA for inter-alpha trypsin inhibitor heavy chain 1, complete cds	0.57	134132	RYANODINE RECEPTOR, SKELETAL MUSCLE	6.9	
330	D89285	S.cerevisiae	0.51	.54.55			
331	Z48951	chromosome XVI	0.57	4210432	(AJ130783) APC2 protein [Mus musculus] TYROSINE	5.3	
332	X95573	A.thaliana mRNA for salt-tolerance zinc finger protein	0.57	1174828	DECARBOXYLASE 2 4.1.1.25) - parsley >gi 169671 (M96070) tyrosine decarboxylase [Petroselinum	5.2	
	·	Xenopus laevis XL- INCENP (XL- INCENP) mRNA,	0.67	445646	PROBABLE ABC TRANSPORTER ATP- BINDING PROTEIN IN NTRA/RPON 5 REGION (ORF1) Azorhizobium caulinodans > gi 311388 (X69959) ORF1	4.0	
333	U95094	complete cds	0.57	465646	(207737) 010 1		
334	AE001116	Borrelia burgdorferi (section 2 of 70) of the complete genome	0.57 .	2314735	(AE000653) Na+/H+ antiporter (nhaA) [Helicobacter pylori 26695] DNA-DIRECTED RNA	4.0	
335	Z34291	R.norvegicus mRNA for putative chloride channel.	0.57	1350832	POLYMERASE I SECOND LARGEST SUBUNIT (RNA POLYMERASE I SUBUNIT 2) chain RPA2 - Euplotes octocarinatus (SGC9) >gi 578407 octocarinatus]	3.0	

Sylvania (**	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
11.00	Nearest N	Neighbor (Blastin Vs. O	endank)	1.000.00.1.00			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					(Z81063) similar to Actinin-type		
		Homo sapiens A30			actin-binding domain containing		
		Vk germline gene,	į		proteins [Caenorhabditis		
336	D88255	partial cds	0.57	3875983	elegans] (U61955) Similar to kinesin-like	3.0	
				_	protein; coded for by C. elegans		
•					cDNA yk184h5.3; coded for by	ł	
					C. elegans cDNA yk184h5.5;	,	
		ļ			coded for by C. elegans cDNA	1	
					yk13d7.3; coded for by C.		
					elegans cDNA yk13d7.5; coded	j	
		•			for by C. elegans cDNA		
		III comis CLI2	ŀ	· .	yk31e1.5; co >gi 3493541	1	
		Homo sapiens SH3-			(AF057567) kinesin-like protein		
		containing adaptor molecule-1 mRNA,			ZEN-4a [Caenorhabditis	ł	
	4 F0270 ()	3	0.57	1397341	elegans	2.3	
337	AF037261	complete cds	0.57	133.371			
	·	Rattus norvegicus	1	1		1	
1	Ì	prostaglandin F2a	1	!		ł	
	}	receptor regulatory			(AF039656) neuronal tissue-	1	
Í		protein precursor,			enriched acidic protein [Homo		
338	U26595	mRNA, complete cds	0.57	2773160	sapiens	2.3	
					(AE001009) quinone-reactive	}	
		R.norvegicus mRNA			Ni/Fe-hydrogenase B-type	l	
1		for interleukin 4	1		cytochrome subunit (hydC)	1.8	
339	X69903	receptor	0.57	2649193	[Archaeoglobus fulgidus]	1.0	
		S.cerevisiae			(U64846) F47D2.5 gene	1	
1		chromosome XV			product [Caenorhabditis	Ì	
١		reading frame ORF	0.57	1458319	elegansi	1.4	
340	Z74825_	YOL083w Foot-and-mouth	0.57	1430313	- Cicquis j		
1		disease virus O vp1	1		proline-rich protein - mouse		
341	AJ131469	gene, strain O/A/58	0.57	91206	(fragment) musculus]	1.4	
341	W131409	Ecirc, Sumii Orroso	1	· · · · · · · · · · · · · · · · · · ·			
1		Mus musculus	1			1	
1		regulator of G-protein		1	1	1	
	i .	signaling 7 (RGS7)				1	
342	AF011360	mRNA. complete cds	0.57	542514	gelsolin - American lobster	0.80	
1	ł	Mus musculus	1		- I - I'- A - origon lobates		
	1	regulator of G-protein	1		gelsolin - American lobster >gi 452313 gelsolin (Homarus		
		signaling 7 (RGS7)	0.57	1078046	>g1 452313 gelsolin (riomatus	0.80	
343	AF011360	mRNA, complete cds	0.57	1078946	jamericanusj	0.00	

	Nearest N	leighbor (BlastN vs. Ge	nbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ			P VALUE	ACCESSION	DESCRIPTION	P VALUE
Ð	ACCESSION	DESCRIPTION	FVALUE	ACCESSION		
		Homo sapiens inosine				
		monophosphate	j		(X77466) 98.8kD polyprotein	
		dehydrogenase type II	1		(Strawberry latent ringspot	0.70
344	L39210	gene, complete cds	0.57	559526	virus]	0.79
		Human endometrial			1	
		bleeding associated			(TO STORE AND STATE OF THE STAT	
	1	factor mRNA,			(K01702) HMW/LMW collagen	0.79
345	U81523	complete cds	0.57	211499	subunit precursor [Gallus gallus]	<u> </u>
						٠.
		Tetrahymena		•	HYPOTHETICAL 100.5 KD]
	1	thermophila			PROTEIN IN IAP-CYSH	[
	1	polyubiquitin (TTU3)			INTERGENIC REGION	
	į	gene, complete cds.			>gi 882654 (U29579) alternate	
	1	and RNA polymerase			gene name ygcB; ORF_f888	1
		II subunit 2 (RPB2)	0.57	2506493	Escherichia colil >gil1789119	0.60
346	U46561	gene, partial cds	0.57	2300.52	NEUROFILAMENT TRIPLET	
			}	1	M PROTEIN (160 KD	1
	1		}	1	NEUROFILAMENT	1
					PROTEIN) (NF-M)	ł
	·	C.japonica mRNA for	ļ	,	>gi 1083164 pir S55395	1
		legumin (clone	1		neurofilament protein M - rabbit	0.46
347	X95543	CiLeg31)	0.57	1709261	(fragment) >gi 854353	0.46
347	1,733,13					
		Homo sapiens mRNA			(AF055904) unknown	0.45
348	Y17282	for cytokeratin type II	0.57	3044086	[Myxococcus xanthus] (AF079369) transcriptional	1
		Frog mRNA fragment	t	1	repressor TUP1 [Dictyostelium	
		for alpha-A2-		242454	discoideum]	0.20
349	X00716	crystallin	0.57	3406654	discoidediti	——
		Klebsiella sp.	1			1
	1	bacteriophage K11	1	1 .		1
	ľ	gene 1 for RNA	0.57	1228093	(Z46913) polyketide synthase	0.16
350	X53238	polymerase	1 0.57	1220073	(S78897) GOR=antigenic	
ĺ	1	H. sapiens FUS gene,			epitope (chimpanzees, Peptide,	
۱ ۵۰۰	200013	exon 12	0.57	# 243898	427 aa] [Pan]	0.090
35	X99012	Human DNA	 	1		
1	1	sequence from PAC	1	1		1
1	1 .	390N22 on	}	1	(U53585) fibronectin attachmer	II 0.053
35:	2 AL008711		0.57	1469545	protein [Mycobacterium avium	0.053
133	712000.11				(U58748) similar to potential	
	1	SOX9 [human, fetal	1		transmembrane domains in S.	
	1	brain, Genomic, 1494		· ·	cerevisiae nulcear division	0.017
1	3 S74506	nt. segment 3 of 3]	0.57	1326350	RFT1 protein (SP:P38206)	0.017

				Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	Nearest N	leighbor (BlastN vs. Go	nbank)	Nearest Neighl	ACIIIS) .		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Human mRNA for		- 	(AF102575) cell surface protein		
		golgi antigen gcp372,			DTFA (Dictyostelium		
354	D25542	complete cds	0.57	4063399	discoideum]	0.005	
		Mus musculus mRNA					
		for alpha 1,3-			Tanagas Paris		
		fucosyltransferase IX,			(Y15732) DNA polymerase beta	7e-11	
355	AB015426	complete cds	0.57	2661842	[Xenopus laevis]	76-11	
		Xenopus mRNA for				}	
		APEG protein.			(Y12090) putative 3,4-	}	
		containing a highly			dihydroxy-2-butanone kinase		
		repetitive amino acid	057	1929056	[Lycopersicon esculentum]	9e-12	
356	X51394	sequence	0.57	1929030	EPIDERMAL GROWTH		
			1		FACTOR RECEPTOR		
					KINASE SUBSTRATE EPS8		
1		Homo sapiens mRNA	ŀ		>gi 530823 (U12535) epidermal		
	Ì	for KIAA0449			growth factor receptor kinase		
257	AB007918	protein, partial cds	0.57	2833239	substrate [Homo sapiens]	3e-13	
357	AB00/918	Homo sapiens mRNA					
İ		for Efs1, complete			(D45027) 25 kDa trypsin	ł	
358	AB001466	cds	0.57	2943716	inhibitor [Homo sapiens]	2e-14	
		Rabbit mRNA for					
1		adult fast skeletal	1		(AC002400) Glutamyl tRNA	2- 20	
359	Y00760	troponin-C	0.57	2576348	synthetase [Homo sapiens]	2e-28	
		H.sapiens brca2 gene	1		(AC004982) similar to yeast	1	
1		exon 3 > ::			hypothetical protein ybk4;	1	
l		emb A62778 A62778			similar to P38164		
		Sequence 19 from	0.67	2410047	(PID: g586461) [Homo sapiens]	2e-55	
360	X95153	Patent WO9719110	0.57	3419847	(FID. 2380401) [Homo suprems]		
١		B.vulgaris mRNA for	0.56	<none></none>	<none></none>	<none></none>	
361	X85967	betavulgin Mycoplasma	0.30	CHOILE			
	1	genitalium DNA				i '	
1	1	gyrase subunit B				1	
	Į.	complete cds, DNA				1	
	l .	polymerase III beta	Ì				
	1	subunit (dnaN) and	1	}		1	
1	i .	seryl-tRNA	.}				
		synthetase (serS)					
362	U09251	genes, partial cds.	0.56	<none></none>	<none></none>	<none></none>	
	1					ł	
	1	Chloroplast Euglena	1	1			
	1	gracilis genes coding	I	1		1	
1	1	for transfer RNAs	1	1	1	i	
	1	specific for threonine	1	1	1		
	.,,,,,,,	glycine, methionine,	0.56	<none></none>	<none></none>	<none></none>	
363	V00158	serine and glutamine.	1 0.36	GIONES			

107	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	racarest I	TEIGHOUT (DIASA VS. O	choank)	rearest recignoof (Diasex 13, 1401-1404-1411)			
SEQ ID	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Clostridium					
		perfringens DNA for					
		D-alanine:D-alanine		•		1	
		ligase, cortical					
		fragment-lytic		\\\C\\\\\\\	NOVE	<none></none>	
364	D88151	enzyme	0.36	<none></none>	<none></none>	<none></none>	
		Methanococcus				1 1	
		jannaschii section 20 of 150 of the					
265	11/7470	complete genome	0.56	<none></none>	<none></none>	<none></none>	
365	U67478	complete genome	0.50	410112	110112	410.12	
		Tachyglossus					
		aculeatus beta-globin		!			
		homolog (HBB)					
366	L23800	gene, complete cds	0.56	<none></none>	<none></none>	<none></none>	
					·		
		Homo sapiens mRNA			·		
		for KIA:A0557				NONT	
367	AB011129	protein, partial cds	0.56	<none></none>	<none></none>	<none></none>	
		Homo sapiens					
		(subclone 10_e10					
368	L77034	from P1 H16) DNA	0.56	<none></none>	<none></none>	<none></none>	
308	L77034	sequence. C.albicans gene for	0.50	CHOILE	110112		
		TFIIIB (BRF1)					
369	Z47202	subunit.	0.56	<none></none>	<none></none>	<none></none>	
		Clostridium					
1		acetobutylicum					
		mannitol-specific					
! !		phosphotransferase		l			
		system (PTS) system,					
		mtlA, mtlR, mtlF, and		•			
,,,	1153060	mtlD genes, complete	0.56	· <none></none>	<none></none>	<none></none>	
370	U53868	cds	0.0	CHONES	1 010		
		Homo sapiens breast		ı			
		cancer putative		I			
		transcription factor					
	/	(ZABCI) mRNA,			1]	
371	AF041259	complete cds	0.56	<none></none>	<none></none>	<none></none>	
		D1					
		Plasmodium					
		falciparum variant-					
		specific surface protein (var-7)					
372	1.42636	mRNA, complete cds.	0.56	2213557	(Z97052) hypothetical protein	8.8	
3/2	L42636	intrava, complete cas.	0.50	±±1333/	(1277032) Hypometical protein		

100	Nearest 1	Veighbor (BlastN vs. G	enbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant Pr	oteins)
SEO						
Ð	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Human protein				
		tyrosine phosphatase	i		THIOREDOXIN REDUCTASE	
		(TEPI) mRNA,			thioredoxin reductase (NADPH)	
373	U96180	complete cds	0.56	731016	[Coxiella burnetii]	8.7
		Homo sapiens PTS			(Y12225) Spi-1/PU.1	
374	L76259	gene, complete cds	0.56	2369863	transcription factor	6.7
		Mus musculus	1			
		D16Jhul7 YAC			hypothetical protein - common	
		98B3 acentric end,			sunflower protein [Helianthus	
375	AF045946	partial sequence	0.56	2130017	annuus]	5.1
		M.musculus mRNA		•	(A COOSDOC) has a stantist	
1	*******	for desmocollin type	0.54	4000001	(AC005936) hypothetical	
376	X97986	1	0.56	4038031	protein [Arabidopsis thaliana]	3.9
1		i	1		COMPONENT SPC42 yeast	
1 1					(Saccharomyces cerevisiae)	
1 1			ŀ		>gi 486054 (Z28042) ORF	
1 1	,		i		YKL042w [Saccharomyces	
					cerevisiae] >gi 666098	
1 1		M.musculus whey	1		(X71621) hypothetical 42.3 kD	
1 1		acidic protein (WAP)	1		protein [Saccharomyces	
377	X79437	gene, exon 1	0.56	549670	cerevisiael	3.9
1	2177437	gene, exon 1		347070	CCIC VISILE)	
1 1		Rat cardiac specific	I		ENDOGLUCANASE G	
1		sodium channel alpha-	- 1		PRECURSOR 3.2.1) CelCCG	
		subunit mRNA.			precursor - Clostridium	
378	M27902	complete cds.	0.56	585234	cellulolyticum cellulolyticum]	3.9
					gp70=envelope protein	
, I	i	Caenorhabditis	j		{endogenous provirus} host=cat	
		elegans cosmid]		lymphoid tissues, Peptide, 445	
379	AF036696	F15B10	0.56	546071	aa]	3.6
					(U14101) putative reverse	
}		ļ	j	•	transcriptase; ORF2; encodes aa	j
1 1			Ţ		motifs conserved in reverse	
) I	-	Caenorhabditis	1		transcriptases: most closely	l
[.]		elegans cosmid	1		related reverse transcriptases are	
1		B0331, complete	-		those of non-LTR	
i		sequence	j		retrotransposons. The 3' 901 bp	}
		[Caenorhabditis	- a - a - b		of this CDS are identical to the	
380	Z99102	elegans]	0.56	603664	3' 901 bp	3.0
	į	Carrie askallina (alama	į.			
] i	1	Equus caballus (clone	j		1	- 1
20,		T131) T-cell receptor	0.56	1070150	because for the factor of the	1.7
381	L27850	DNA. V-region.	0.30	1079150	transcription factor shn - fruit fly	1.7

23.5	Nonzacz N	Neighbor (BlastN vs. Go	nhank) I	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	Nearest N	reignbor (Diasus vs. Of	ilivalik)	remest reign	1		
SEQ	A CCECCION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
10	ACCESSION	DESCRIPTION	1 VALUE	ACCESSION.	HYPOTHETICAL 113.1 KD		
		M.musculus mRNA			PROTEIN IN PRE5-FET4		
			į		INTERGENIC REGION		
200	*******	for desmocollin type	0.56	2497227	>gi 1072409 (Z54141) unknown	1.7	
382	X97986	Didelphis virginiana	0.30	2491221	(U12964) contains ankyrin-like		
		G protein receptor	į		repeats; similar to human		
•		kinase 2 mRNA.	, ,		desmoplakin repeat region		
202	A 5007466		0.56	1213453	[Caenorhabditis elegans]	1.3	
383	AF087455	complete cds Human mRNA for	0.30	1215455	(Cachornacanas siegens)		
1			ŀ		\	1	
	500011	KIAA0189 gene,	0.56	226535	protease [Hepatitis B virus]	1.1	
384	D80011	complete cds	0.30	220333	processe (riepanus 2 va ss)		
		Mus musculus mRNA	·		(AB014572) KIAA0672 protein		
322	. 100000	for HAP1-A protein,	0.56	3327158	[Homo sapiens]	1.0	
385	AJ002272	3' region	0.36	3321138	(LIOINO SAPICIIS)	<u>-</u> -	
l	1		•				
l		TT				j	
1		Homo sapiens inosine	1				
l		monophosphate	į		coat protein - strawberry latent]	
		dehydrogenase type II	0.56	628431	ringspot virus	0.77	
386	L39210	gene, complete cds	0.36	020431	Thigspot virus	0,,,,	
l		Mouse Thy-1.2 gene			(AB014516) KIAA0616 protein		
205	,,,,,,,,,,	5' untranslated region	0.56	3327046	[Homo sapiens]	0.59	
387	X02770	and exon 1	0.30	3327040	(Homo suprens)	- 0.02	
1	!	Schizosaccharomyces			ŀ		
.	1	pombe Wiskott-					
}	1	Aldrich Syndrome			salivary proline-rich		
ļ	ŀ	protein homolog			phosphoprotein precursor PRH1	1	
1	Į .	(wspl+) genc,			(allele PIF) - human >gi 190484		
t		complete cds, and			(K03203) prepro salivary		
	1	BTF3/beta-NAC	ļ ,		proline-rich protein [Homo	(
200	A E039636	gene, partial sequence	0.56	88466	sapiens] >gi 190512	0.35	
388	AF038575	Rat mRNA for fetal	0.50	00700			
		intestinal lactase-					
ľ		phlorizin hydrolase	1		(Z48674) chitinase homologue		
389	X56747	precursor, partial	0.56	2072742	[Sesbania rostrata]	0.23	
307	A30/4/	Garboreum mRNA					
		for farnesyl				(
1	!	pyrophosphate			(X07882) Po protein [Homo	j i	
390	Y12072	synthase	0.56	296670	sapiens]	0.20	
1750	112072	Symulase					
	{	p15=cyclin D-			1		
	1	dependent kinases 4				!	
		and 6-binding					
		protein/p15 product			protein kinase (EC 2.7.1.37)]	
	1	[exon/intron 1]			SPRK - human sapiens]		
		human, brain tumors,			>gi 1090771 prf 2019437A		
391	S75756	Genomic, 753 nt]	0.56	1082743	protein Tyr kinase I	0.15·	
721	3/3/30	Cellotine, 735 mj	<u> </u>				

- Marin	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neigh	nbor (BlastX vs. Non-Redundant P	roteins)
SEQ				i demest i del Bu	1001 (Blasca vs. 14011-Redulidant P	(Silis)
ID ID	ACCESSION	DESCRIPTION	DVALLE	+ COECCIO:	DESCRIPTION:	
-	ACCESSION		PVALUE	ACCESSION	DESCRIPTION	P VALUE
		Equus caballus type				
392	1162620	11 collagen mRNA.	0.56		[Segment 1 of 2] COLLAGEN	
392	U62528	complete cds C.reinhardtii mRNA	0.56	461671	ALPHA 1(I) CHAIN	0.030
		for unknown lumenal	1		(1003672)	
393	X96877	polypeptide	0.54	22/1670	(AC003672) putative zinc finger	1
	A700//	porypeptide	0.56	3341678	protein [Arabidopsis thaliana]	5e-09
]	!		1			
			1	1		[
					(AL009196) 1-	
					evidence=predicted by content;	
		٠,			1-method=genefinder;084; 1-	
				·	method_score=59.41; 1-	•
					evidence_end; 2-] .
	,				evidence=predicted by match; 2-	
			:		match_accession=AA950019; 2-	İ
	<u> </u>	cGATA-3 [chickens,			match_description=LD29959.5p	
		liver, Genomic, 979			rime LD Drosophila	
394	S78788	nt, segment 4 of 4)	0.56	2661590	melanogas	2e-11
					(U41534) coded for by C.	
		Drosophila			elegans cDNA CEES142F;	
		melanogaster Ste20-	<u> </u>		Similar to helicases of	
205	A FOOSS AG	like protein kinase	[•	SNF2/RAD54 family.	
395	AF006640	mRNA, complete cds	0.56	1109830	[Caenorhabditis elegans]	6e-12
		Drosophila		,	(U41534) coded for by C.	
ļ		melanogaster Ste20-			elegans cDNA CEESI42F; Similar to helicases of	
l		like protein kinase	ı		SNF2/RAD54 family.	j
396		mRNA, complete cds	0.56	1109830	[Caenorhabditis elegans]	4e-13
	. = 0000,0		0.30	1102030	(AL030996) dJ1189B24.4	46-13
			i		(novel PUTATIVE protein	
İ	j	Ì	1		similar to hypothetical proteins	ľ
ł		Aquifex aeolicus	Į		S. pombe C22F3.14C and C.	ľ
ŀ		section 48 of 109 of	1	••	elegans C16A3.8) [Homo	
397		the complete genome	0.56	3688350	sapiens)	3e-66
		S.cerevisiae				
		chromosome II	I			
		reading frame ORF				- 1
398		YBR210w	0.55	<none></none>	<none></none>	<none></none>
		Mus musculus mRNA				
,,,	1	for ubiquitin				
399		onjugating enzyme	0.55	<none></none>	<none></none>	<none></none>
- 1		Homo sapiens	- 1			
		subclone 2_g5 from BAC H107) DNA	•			
400	1	· I	0.55	ANONE	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	NOVE
700	AC001-01	еqueпсе	0.55	<none></none>	<none></none>	<none></none>

is possi	Negros	Neighber (Blook) C	· ·	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
		Neighbor (BlastN vs. C	renoank)	Nearest Neig	nbor (BlastX vs. Non-Redundant P	oteins)	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Alouatta seniculus					
]		breast and ovarian					
1		susceptibility	Ì		1	ł	
	1	(BRCA1) gene,		ŀ			
401	AF019079	partial cds	0.55	<none></none>	<none></none>	<none></none>	
		Human serglycin					
1		gene, exons 1,2, and		i		I	
402	M90058	3.	0.55	<none></none>	<none></none>	<none></none>	
403	AB013469	Mus musculus CLM2 gene for cytohesin 2, complete and partial cds, alternative splicing	0.55	1729760	(Z68152) chitinase [Gossypium	8.6	
			0.55	1723700	misdani	8.0	
·					PHOTOSYSTEM II 10 KD PHOSPHOPROTEIN deltoides] >gi 2143326 gnl PID e319090 (Y13328) 10kDa		
		Bacteriophage P1 ban			phosphoprotein [Populus	I	
404	AJ011592	gene	0.55	2493689	deltoides]	6.6	
		T.brucei kinetoplast			(AF049132) NADH		
105	5.5. 6.6	maxicircle variable	1		dehydrogenase subunit 5		
405	Z15118	region DNA	0.55	2970432	[Florometra serratissima]	6.5	
		S.cerevisiae	ŀ				
406	Z48951	chromosome XVI cosmid 9723	0.55	4010400	(AJ130783) APC2 protein [Mus		
700	240951	Homo sapiens mad	. 0.33	4210432	musculus)	4.9	
407		protein homolog Smad2 gene, promoter, exon la and exon lb	0.55	3319290	(AF055994) thyroid hormone receptor-associated protein complex component TRAP220 [Homo sapiens]	4.9	
					L		
408		Homo sapiens genomic DNA, 21q region, clone: PH11Bm42	0.55	125694	KRUEPPEL PROTEIN >gi 72899 pir TWFF Krueppel gap protein - fruit fly (Drosophila sp.) melanogaster] >gi 224875 prf 1202348A	2.8	
		/111 IDIII72	0.55	125684	X-LINKED PEST-	3.8	
J			-		CONTAINING	ľ	
	·	Plasmodium vivax	j		TRANSPORTER transporter -		
		najor blood stage			human >gi 458255 (U05321) X-	f	
- 1		urface antigen gene.			linked PEST-containing		
409		partial cds.	0.55	549453	transporter [Homo sapiens]	3.8	
				0.5.00	mansporter (Fronto suprens)	٥.٥	

SEQ ID ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION P V ID ACCESSION DESCRIPTION DESCRIPTION P V ID ACCESSION DESCRIPTION DESCRIPTION P V ID ACCESSION DESCRIPTION DESCRIPTION DESCRIPT	東震震	Negroo	National Property			· .	
D ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION P V			Theighbor (Blastin vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant P	roteins)
Fugu rubripes mRNA for sodium channel alpha subunit, partial cds 0.55 1435038 (D38024) ORF [Homo sapiens] 3 3 3 3 3 3 3 3 3	-	1					
Fugu rubripes mRNA for sodium channel alpha subunit, partial cds	_ <u>m</u>	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
Application			Fugu rubripes mRNA	4			I VALUE
10 D37977 cds Ostertagia ostertagic cathepsin B-like Cysteine protease gene, partial cds. O.55 3941277 Ostertagia ostertagic cathepsin B-like Cysteine protease gene, partial cds. O.55 3941277 Ostertagia ostertagic cathepsin B-like Cysteine protease gene, partial cds. O.55 3941277 Ostertagia ostertagic mitotic phosphoprotein 44 Cysteine protein 44 Dysteine protein 44 Cysteine protein 44 Cysteine protein 44 Cysteine protein 44 Cysteine protein 44 Cysteine protein 65 Cysteine protein 67 Cysteine protein 7 Cysteine protein 8 Cysteine protein 9 Cysteine 9 Cysteine 9 Cysteine protein 9 Cysteine 9 Cy	1 1		for sodium channel				
Ostertagia ostertagic cathepsin B-like cysteine protease gene. partial cds. 0.55 3941277 (AF000900) p45 [Rattus norvegicus] 2 2 2 2 2 2 2 2 2		l	alpha subunit, partial	1			
Ostertagia ostertagic cathepsin B-like cysteine protease gene, partial cds. 0.55 3941277 (AF000900) p45 [Rattus norvegicus] 2 2 2 2 2 2 2 2 2	410	D37977		0.55	1435038	(D38024) ORE (Homo conional	1
Cysteine protease gene, partial cds. 0.55 3941277 (AF000900) p45 [Rattus norvegicus] 2						(220024) Old [Homo sapiens]	3.7
M88505 gene, partial cds D.55 3941277	i l	j .			1		l .
Macsay Refere Partial cds Ca	1 1					(AF000900) p45 (Rame	1
Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds 0.55 2570154 PP1 [Sus scrofa] 2 2 2 2 2 2 2 2 2	411	M88505		0.55	3941277		2.9
1.2 U95098 Phosphoprotein 44 mRNA, partial cds U89241 Human mibp gene, partial cds U89241 Partial cds U89241 Exercise U89241	1		Xenopus laevis				2.9
Digital Phosphoprotein 44 mRNA, partial cds 0.55 2570154 potentiated inhibitory protein of PP1 [Sus scrofa] 2				1	1	(AB008376) 17-kDa PKC-	
Human mibp gene, partial cds 0.55 2570154 PP1 [Sus scrofa] 2				1	İ		i i
Human mibp gene, partial cds	412	U95098.		0.55	. 2570154		2.8
AF006821 Partial cds Sequence Sequen							2.6
Survival of motor neuron protein interacting protein I (SIP1) mRNA. 414 AF027151 complete cds 0.55 4007790 protein [Schizosaccharomyces pombe] 415 AF006821 partial cds 0.55 2245075 protein 1 416 Y12736 and orfX 0.55 3386334 familiaris] 417 U38307 sequence. 0.55 1362802 (fragment) >gil547517 1.3 Mouse mRNA for Rad51 protein 0.55 1374698 NP220 (MPRO expirited).	413	U89241	partial cds	0.55	4097465		2.2
neuron protein	ŀ			1			
interacting protein I (SIP1) mRNA. complete cds Bufo marinus natriuretic peptide receptor C mRNA. partial cds Lactococcus lactis cremoris plasmid pJW565 DNA. llabiiM, llabiiR genes and orfX Mus musculus collagen alpha-1 type 1 gene. 5' flanking region, partial 417 U38307 Mouse mRNA for Mouse mRNA for And AF027151 Interacting protein I (SIP1) mRNA. AF007790 A00779	1			1		- 1	
AF027151 Complete cds 0.55 4007790 Strand polynucleotide binding protein [Schizosaccharomyces pombe] 1.	I			i		(AL034463) putative single-	
AF027151 Complete cds 0.55 4007790 protein [Schizosaccharomyces pombe] 1.	Į					strand polynucleotide binding	
AF027131 Complete cds 0.55 4007790	414	4 E027161				protein [Schizosaccharomyces	j
natriuretic peptide receptor C mRNA, partial cds Lactococcus lactis cremoris plasmid pJW565 DNA, llabiiM, llabiiR genes and orfX Mus musculus collagen alpha-1 type 1 gene. 5' flanking region, partial sequence. 1.3 Mouse mRNA for Rad51 protein 1.5 (Z97343) GTP-binding RAB2A protein (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (D83032) nuclear protein, partial protein, processing and processing all protein, processing and pro	414	AF02/151	Complete cds	0.55	4007790		1.7
AF006821 receptor C mRNA, partial cds 0.55 2245075 protein 1.7				1			
AF006821 partial cds 0.55 2245075 protein 1.7							
Lactococcus lactis cremoris plasmid pJW565 DNA, llabiiM, llabiiR genes and orfX Mus musculus collagen alpha-1 type 1 gene. 5' flanking region, partial sequence. 0.55 0.55 1362802 (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3 (Dasios) 1.3	415	AF006821				(Z97343) GTP-binding RAB2A	i
cremoris plasmid pJW565 DNA. llabiiM, llabiiR genes and orfX Mus musculus collagen alpha-1 type 1 gene. 5' flanking region, partial sequence. 0.55 1362802 (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (AF035120) type I procollagen pro-alpha 2 chain [Canis familiaris] 1.3 (Bassing mucin - human (Gragment) >gi 547517 1.3 (D83032) nuclear protein, Rad51 protein 1.3 (D83032) nuclear protein, Rad51 protein	+	11 000021		0.55	2245075	protein	1.7
PJW565 DNA,	ı						
AlfabiiM, IlabiiR genes and orfX 0.55 3386334 pro-alpha 2 chain [Canis familiaris] 1.3						/47005100	
416 Y12736 and orfX 0.55 3386334 familiaris] 1.3	1					(AF035120) type I procollagen	
Mus musculus	416			0.55	2206224		j
1 gene. 5' flanking gastric mucin - human				0.55	3380334	[Tamiliaris]	1.3
1 gene. 5' flanking gastric mucin - human	l'	.	collagen alpha-1 type	1			i
417 U38307 sequence. 0.55 1362802 (fragment) >gi 547517 1.3				1		1	1
417 U38307	ł	Į:	region, partial	. 1		eastric mucin, human	- 1
Mouse mRNA for (D83032) nuclear protein,	417	U38307	sequence.	0.55	1362802		., 1
416 D134/3 Rad5 protein 0.55 1374698 NP220 (Home against	- 1	[1	Mouse mRNA for				1.3
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						The provide sapiens	1.3
acetylcholinesterase	1					[i
gene, alternatively	ı			- 1		1	
spliced products. (Z94752) hypothetical protein				1		(Z94752) hypothetical protein	1
419 Ar043238 partial cds 0.55 3261734 RV1004c	119			0.55	3261734		0.99
Methanobacterium	- 1						
thermoautotrophicum	1			1		1	1
from bases 1 to	- 1			j			
10208 (section 1 of	j			1			1
420 AE000795 genome 0.55 186306 (M94131) mucin [Homo	20 .					(M94131) mucin [Homo	1
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7	Neare	st Neighbor (BlastN vs.	Genhank)	Nonros Nois	hts Ol W		
SE			Jenoank)	Nearest Neig	ghbor (BlastX vs. Non-Redundant	Proteins)	
ID	~	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	D	
		Y.lipolytica SEC62	T		(Z81068) F25H5.2	P VALU	<u>'</u>
42	X99537	депе	0.55	3876397	[Caenorhabditis elegans]	0.50	_
	1	Aquilegia sp.			(Cachernabarus elegans)	0.58	_
- 1	1	phytochrome	ļ			1	
1 422	T. T. T. T. T. T. T. T. T. T. T. T. T. T	(PHYB/D) gene,			(AF005370) ribonucleotide-	1	
422	U08147	partial cds.	0.55	2338024	reductase, large subunit	0.57	
1	ł	Warrier G C DVA					-
ļ	1	H.sapiens CpG DNA	•	-		ı	
423	Z56586	clone 12c8, reverse read cpg 12c8.rt1d.	0.55		(U46007) espin [Rattus		i
	250500	read cpg12co.ft1d.	0.55	3320122	norvegicus]	0.44	ı
1.	1	Mus musculus	1				٦
1	1	glutamine:fructose-6-	.	•	T.		
1		phosphate				ł	ı
1	1	amidotransferase			1	1	ı
1 -	1	(GFAT) gene, 5'	ļ (hypothetical protein -	1	ı
424	U39442	region and partial cds	0.55	282600	Mycoplasma hyorhinis	0.43	1
		Rat chymotrypsin B			in jeopiasina nyoniniis	0.43	4
1	Į.	(chyB) gene,	1 1		(Y17034) Bassoon [Mus	ł	ı
425	K02298	complete cds.	0.55	3413810	musculus]	0.33	ı
120	V0.1707	M.musculus clusterin				0.55	ł
426	X84792	gene	0.55	1652475	(D90905) hypothetical protein	0.25	ı
		Capra aegagrus Saanen and Weisse	1				1
1		Edel breeds DR beta-				1	l
ı	ĺ	chain antigen binding	1			1	l
1		domain, MHC class II	i i		Gringer to the second		ĺ
427	U00185	DRB	0.55	2507136	SUBTILIN BIOSYNTHESIS	1	l
				2507130	PROTEIN SPAB	0.19	l
1 1		H. sapiens CpG DNA,			l	1	l
		clone 178a12, reverse			(M17294) unknown protein		l
428	Z54946	read cpg178a12.rtla.	0.55	807646	[Human herpesvirus 4]	0.065	
						0.005	
		Oryctolagus	1	•		1 1	
		cuniculus anion	1				
		exchanger 3 brain	l				
429	AF031650	isoform (AE3)			(U68412) fibrillar collagen		
 -	100100U	mRNA. complete cds	0.55	1778210	[Arenicola marina]	0.044	
I		Bovine adenylyl	- 1		(4.500000)		
	i	cyclase Type I	1		(AE000997) conserved	•	
430	M25579	mRNA. complete cds.	0.55		hypothetical protein	I	
		H.sapiens Ski-W	- 5.55	2047040	[Archaeoglobus fulgidus]	0.023	
431		mRNA for helicase	0.55	330452	(M14708) DNA polymerase	0.000	
			3.55	220422	[Human cytomegalovirus]	0.023	

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CDNA EST EMBL:C11959 Comes from this gene; cDNA EST EMBL:C10341 comes from this gene; cDNA EST yk494e4.3 comes from this gene; cDNA EST yk494e4.3 comes from this gene; cDNA EST yk494e4.3 comes from this gene; cDNA EST yk494e4.3 comes from this gene; cDNA EST yk494e4.3 comes from this gene; cDNA EST yk494e4.3 comes from this gene; cDNA EST yk494e3.5 comes from this gene [Caenorhabditis elegans] 7e-06				1 1			Į.
comes from this gene; cDNA EST EMBL:C10341 comes from this gene; cDNA EST yk494e4.3 comes from this gene; cDNA EST yk494e4.3 comes from this gene; cDNA EST yk494e3.5 comes from this gene; cDNA EST yk494e3.5 comes from this gene; cDNA EST yk494e3.5 comes from this gene; cDNA EST yk494e3.5 comes from this gene; cDNA EST yk494e3.5 comes from this gene; cDNA EST yk494e3.5 comes from this gene; cDNA EST EMBL:C10341 comes from this gene; cDNA FSO COLOTION FSO EMBL:C10341 comes from this gene; cDNA FSO COLOTION FSO EMBL:C10341 comes from this			}	1 1			i
Human I kappa B epsilon (IkBe) Arabidopsis thaliana Atpk7 gene for serine/threonine protein kinase, complete cds CZ81505) Similarity to Metanococcus hypothetical protein 0682 (TR:Q58095) Swinepox virus complete ORFS C20L-C1L > :: gb I58297 I58297 Sequence 14 from patent US 5651972 Human immunodeficiency virus type 1 env gene EST EMBL:C10341 comes from this gene; cDNA EST yk448a8.5 comes from this gene; cDNA EST yk444aa.5 comes from this gene; cDNA EST yk444aa.5 comes from this gene; cDNA EST yk444aa.5 comes from this gene; cDNA EST yk444aa.5 comes from this gene; cDNA EST yk444aa.5 comes from this gene; cDNA EST yk444aa.5 comes from this gene; cDNA EST yk444aa.5 comes from this gene; cDNA EST yk444aa.5 comes from this gene; cDNA EST yk444aa.5 comes from this g				1 1			- 1
Human I kappa B epsilon (IkBe) mRNA, complete cds 0.55 3875577 (Caenorhabditis elegans) 7e-06				1 1			I
Human I kappa B				1			
Human I kappa B epsilon (IkBe) mRNA, complete cds 0.55 3875577 Gaenorhabditis elegans 7e-06				1 1			1
epsilon (IkBe) mRNA, complete cds Arabidopsis thaliana Atpk7 gene for serine/threonine protein kinase, complete cds Swinepox virus complete ORFS C20L-C1L > :: gb I58297 I58297 Sequence 14 from patent US 5651972 Human immunodeficiency virus type 1 env gene 0.55 3875577 [Caenorhabditis elegans] 7e-06 (Z81505) Similarity to Metanococcus hypothetical protein 0682 (TR: Q58095) [Caenorhabditis elegans] 4e-42 CNONE> NONE> NONE> NONE>			Human I kanna B	i i			1
433 U91616 mRNA, complete cds 0.55 3875577 [Caenorhabditis elegans] 7e-06	- 1			1 1			1
Arabidopsis thaliana Atpk7 gene for serine/threonine protein kinase, complete cds Swinepox virus complete ORFS C20L-C1L > :: gb 158297 158297 Sequence 14 from patent US 5651972 Human immunodeficiency virus type 1 env gene 0.54 C381505) Similarity to Metanococcus hypothetical protein 0682 (TR:Q58095) (Caenorhabditis elegans) 4e-42 Caenorhabditis elegans) 4e-42 Caenorhabditis elegans Atpk7 gene for (Z81505) Similarity to Metanococcus hypothetical protein 0682 (TR:Q58095) (Caenorhabditis elegans) 4e-42 Caenorhabditis elegans 4e-42 Caenorhabditis elegans Caenorhabditis elegans Atpk7 gene for (Z81505) Similarity to Metanococcus hypothetical protein 0682 (TR:Q58095) (Caenorhabditis elegans) 4e-42 Caenorhabditis elegans Caenorhabditis elegans Atpk7 gene for (Z81505) Similarity to Metanococcus hypothetical protein 0682 (TR:Q58095) (Caenorhabditis elegans) Caenorhabditis elegans Atpk7 gene for (Z81505) Similarity to Metanococcus hypothetical protein 0682 (TR:Q58095) (Caenorhabditis elegans) Ac-42	433			0.55	20 76577		
Atpk7 gene for serine/threonine protein kinase, complete cds				- 0.55	. 3013311	[Caenorhabditis elegans]	7e-06
Serine/threonine Protein kinase, Complete cds O.55 3876072 Protein 0682 (TR:Q58095) Gaenorhabditis elegans 4e-42	Į] [(781505) Similaring to	j
Protein kinase, protein 0682 (TR:Q58095)	j]
Swinepox virus Swinepox virus Caenorhabditis elegans 4e-42	ł]	protein kinase,	1			1
Swinepox virus	434	D10910	complete cds	0.55	3876072	1	40-42
C20L-C1L > :: gb I58297 I58297 Sequence 14 from patent US 5651972			Swinepox virus			(Carioniabana ciegana)	46-42
gb I58297 I58297 Sequence 14 from patent US 5651972 0.54 <none> <none> Human immunodeficiency virus type 1 env gene 0.54 <none> <none> <none> <none></none></none></none></none></none></none>	1			1]
Sequence 14 from patent US 5651972 0.54 <none> <none> </none></none>	- 1			j			- 1
135 L22013 patent US 5651972 0.54 <none> <none> </none></none>	-		•		İ		[
Human immunodeficiency virus type 1 env gene 0.54 <none> <none> <none> <none> <none></none></none></none></none></none>	47,			1	ļ	1	1
Human immunodeficiency virus type 1 env gene 0.54 <none> <none> <none></none></none></none>	435	L22013	patent US 5651972	0.54	<none></none>	<none></none>	NONE>
immunodeficiency 436 Z92653 virus type 1 env gene 0.54 <none> <none> <none></none></none></none>		1.					
136 Z92653 virus type env gene 0.54 <none> <none> <none> </none></none></none>				i		. [
ZNONES ZNONES	436						1
	.55	272033	in us type I env gene	0.54		<none></none>	NONE>

5編後	Neares	Neighbor (BlastN vs.	Genhank)	Norman N.	th (D) v	
SEC		Treighbor (Diasus Vs.	Jenbank)	ivearest Neig	ghbor (BlastX vs. Non-Redundant	Proteins)
D	-	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
<u> </u>						
		E.coli phosphate-				+
1		repressible	1 .	i		1
	ı	periplasmic	1 .		1	1
	1	phosphate-binding	1			ł
1	1	protein (phoS),	j	1	į.	
		peripheral membrane	1	ł		
1		proteins (pstC. pstB and phoU) and	l i		ļ	
1	I					
1	1	integral membrane	1		,	1 1
437	K01992	protein (pstA) genes, complete cds.	0.54		·	
1 43/	K01992	Plasmodium	0.54	· <none></none>	<none></none>	<none></none>
1	I	falciparum			}	
1		chromosome 2.]			
	j	section 52 of 73 of	l		İ	1 1
]	ŀ	the complete	1		ì	1 1
438	AE001415	sequence	0.54	<none></none>	<none></none>	<none></none>
					CHORLE	GAONES
!		Helianthus tuberosus	1		İ	1
1		lectin 2 mRNA,	1			j
439	AF064030	complete cds	0.54	<none></none>	<none></none>	<none></none>
1		E.coli plasmid DNA				10112
440	X12591	for colicin E9	0.54	<none></none>	<none></none>	<none></none>
i .		Comment	1			T
i		Caenorhabditis				j i
441	U73679	elegans YNK1-a mRNA, complete cds			1	1 1
H	013019	Unidentified	0.54	<none></none>	<none></none>	<none></none>
1		bacterium DNA for	1		1	1 1
442	Z93990	16S ribosomal RNA	0.54	-NONE-		1
	2,3,7,0	B.vulgaris mRNA for		<none></none>	<none></none>	<none></none>
443	X85967	betavulgin	0.54	757836	(Z37980) ORF12 [Escherichia coli]	
				, , , , , , , ,	[COII]	8.3
		Sambucus nigra	İ			
		ribosome inactivating	ŀ			1
		protein precursor			(M80653) tetraheme	1
444	U76524	mRNA. complete cds	0.54 .	151377	[Pseudomonas stutzeri]	6.2
<u> </u>	J,	U	- 1		1	· [
- [H.sapiens gene for 5S	1			- 1
ŀ		RNA (640 bp) > ::	1			ļ
- 1		mb X71801 HS5SR6	ł]
445		10B H.sapiens gene	054	^	(AE001216) T. pallidum	I
		for 5S rRNA (640 bp) Human mibp gene,	0.54	3322653	predicted coding region TP0369	2.7
446		partial cds	0.54		(U62253) 16kDa secretory	
			4-6.0	4097465	protein [Sus scrofa]	2.2

-1740 X	Nearest	Neighbor (BlastN vs. C	enhanki	Magazza Maia	hhor (BlassV N. B. 1. 1. B.	
SEO	ivearest	TACIBIDOL (DIASUA VS. C	choank)	inearest ineigi	hbor (BlastX vs. Non-Redundant P	roteins)
D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
						†
447	L16013	Rattus norvegicus Q- like gene sequence	0.54	3087760	(AJ005583) p75 protein [Crypthecodinium cohnii]	0.95
448	U60275	Capra hircus skeletal muscle voltage-gated chloride channel gClC-1 mRNA, partial cds	0.54	1781344	(Y10438) FK506 polyketide synthase	0.95
449	U36795	Myxococcus xanthus rfbABC O-antigen biosynthesis operon, rfbA, rfbB, and rfbC genes, complete cds.	0.54	3877232	(Z81540) predicted using	0.74
450	AF053091	Drosophila melanogaster eyelid (eld) mRNA, complete cds	0.54	2144110	zinc finger protein RIZ - rat	0.14
451	V00602	Genome of the bacteriophage fd (Inoviridae).	0.54	2661620	(AL009197) hypothetical	0.11
452		Human semaphorin (CD100) mRNA, complete cds	0.54	125682	KERATIN. ULTRA HIGH- SULFUR MATRIX PROTEIN (UHS KERATIN) >gi 109116 pir A36686 ultra- high-sulfur keratin - sheep >gi 1306 (X55294) ultra high- sulphur keratin protein [Ovis aries]	0.003
453		5.coelicolor secD.	0.54	3874972	(Z99709) similar to Elongation factor Tu family (contains ATP/GTP binding P-loop); cDNA EST EMBL:D76223 comes from this gene; cDNA EST yk478c5.5 comes from this gene [Caenorhabditis elegans]	7e-06
]	H.sapiens mRNA for DAN26 protein,		3014712	(Z70750) similar to vanadate resistance protein transmembranous domains	/6-00
454	Y08265	partial	0.54	3875131	[Caenorhabditis elegans]	5e-12

是於漢		Neighbor (BlastN vs. C	Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant l	Proteins)
SEQ	1					10.01.12)
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU
		Hydromantes				11 VALU
		platycephaius		<u> </u>		-
	l	cytochrome b (cytb)		Ĭ		ì
	Į.	gene, mitochondrial	1		·	1
	· /	gene encoding	l	[j
		mitochondrial	i	1		-
455	U89613	protein, partial cds	0.53	<none></none>	<none></none>	<none:< td=""></none:<>
		Habrobracon hebetor				1
		cytochrome oxidase			1	
		II gene, partial cds;				
		and tRNA-Asp, tRNA				
		His, and tRNA-Lys		•	ł	.
		genes, complete				i
ı		sequence,				1
		mitochondrial genes				1
1		for mitochondrial				
456	AF034597	products	0.53	<none></none>	<none></none>	<none></none>
ı						CHOINES
ľ		Yeast (S.cerevisiae)	I		1	1
457		tau repetitive element			1	ļ
43/		and Cys-tRNA. Human mRNA for	0.53	<none></none>	<none></none>	<none></none>
I						
458		actin-binding protein	0.53		bullous pemphigoid antigen 2 -	
-	N23410	(maini)	0.53	2134839	human	6.2
ŀ	1		. [
- 1		Drosophila	1			
- 1		subobscura alchohol	İ		1	
ı		dehydrogenase (Adh)	i			
- 1	[8	gene, and alchohol	į.		1	
1		lehydrogenase (Adh-	- 1	•	1	
ا دم		lup) gene, complete	1		hair keratin cysteine rich protein	
59	M55545	ds's.	0.53	2136865	- sheep	2.1

- BA778	No	Valuables (Dissaline C	anhank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
河北	Nearest I	Veighbor (BlastN vs. G	епрапк)	Mearest Neight	l (Diasix vs. Non-Redundant Pr	otems) .	
SEQ					procession.		
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
		Methanobaeterium					
		thermoautotrophicum					
		methylene-					
}		tetrahydromethanopte		•	HYPOTHETICAL 91.6 KD		
		rin dehydrogenase			PROTEIN IN HXT8-CRT1		
		(mtd),			INTERGENIC REGION		
		imidazoleglycerol-			>gi 1078261 pir S50773		
		phosphate			probable membrane protein		
1		dehydrogenase			YJL212c - yeast		
		(hisB), and putative			(Saccharomyces cerevisiae)		
	. (ferredoxin (fdxA)			>gi 496950 (Z34098) ORF [Saccharomyces cerevisiae]		
		genes, complete cds, orf9 gene, partial cds,			>gi 1015596 (Z49487) ORF		
460	U19362	orfs	0.53	731969	УЛL212c	0.54	
400	019302	0115	0.55	731909	11522120	0.54	
					GERM CELL-LESS PROTEIN	1	
					fruit fly (Drosophila		
		Rattus norvegicus			melanogaster) >gi 157490		
1		mRNA for MEGF1,			(M97933) germ cell-less protein		
461	AB011527	complete cds	0.53	417037	[Drosophila melanogaster]	3e-06	
		Darillan Samua Mary D	j				
ł		Bacillus firmus MsyB gene, 5' upstream					
462	U64313	region and partial cds	0.52	<none></none>	<none></none>	<none></none>	
102	004313	Caenorhabditis		41.01.125			
		elegans paraquat	i		1		
		responsive protein	i				
		(CePqM132) mRNA,					
463	AF008590	complete cds	0.52	<none></none>	<none></none>	<none></none>	
		Mus saxicola					
		spermidine/spermine	1	•			
		N1-acetyltransferase		•			
	•	(SSAT) gene.	ľ				
464	L10245	complete cds.	0.52	<none></none>	<none></none>	<none></none>	
					INSULIN-LIKE GROWTH		
į į					FACTOR IB PRECURSOR		
		Arabidopsis thaliana			(IGF-IB) (SOMATOMEDIN C)		
Į į	i	cellulose synthase			>gi 69361 pir IGHU1B insulin-		
		catalytic subunit (Ath-	. 1		like growth factor IB precursor - human prepropeptide [Homo		
465	AF027173	A) mRNA, complete cds	0.52	124263		7.7	
05	TT-0-1113	Cus	U.J4	127203	lanhiena]	··/	

WO 01/02568 PCT/US00/18374

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
機器	Nearest N	Veighbor (BlastN vs. G	nbank)	Nearest Heighbor (Shaber 19, 1991)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Caenorhabditis					
		elegans cosmid					
		H31B20, complete			1		
		sequence					
		[Caenorhabditis			(D88451) aldehyde oxidase [Zea		
466	AL021066	elegans	0.52	2589162	mays]	6.0	
+00	ALUZIOU	·			(U39850) coded for by C.		
		·			elegans cDNA yk37g1.5; coded		
					for by C. elegans cDNA		
			1		yk5c9.5; coded for by C.		
		Porphyra linearis 18S			elegans cDNA ykla9.5;		
		ribosomal RNA gene,			alternatively spliced form of		
467	AF038588	3' partial sequence	0.52	1055055	F52C9.8b	4.6	
-40/	VI 020200	J partie sequence					
		Borrelia burgdorferi			· ·		
i		(section 11 of 70) of			(AB021287) polyprotein		
468	AE001125	the complete genome	0.52	4115827	[Hepatitis G virus]	2.0	
400	AE001123	uic complete general			(U41534) coded for by C.		
		Drosophila	į į		elegans cDNA CEESI42F;		
		melanogaster Ste20-			Similar to helicases of		
		like protein kinase]		SNF2/RAD54 family.		
469	AF006640	mRNA. complete cds	0.52	1109830	[Caenorhabditis elegans]	0.002	
100	711 0000-10	Aplysia californica					
1		ubiquitin carboxyl-			· ·		
		terminal hydrolase	<u> </u>			1	
1		(Ap-uch) mRNA,					
470	U90177	complete cds	0.51	<none></none>	<none></none>	<none></none>	
<u> </u>		S.cerevisiae				ł	
	ļ.	chromosome XI				į.	
1		reading frame ORF					
471	Z28304	YKR079c	0.51	<none></none>	<none></none>	<none></none>	
		Caenorhabditis					
1		elegans cosmid			1		
1		R03E1, complete	1	. '	1		
i		sequence				1	
1		[Caenorhabditis		1	HYDROPHOBIC SEED	1 7/	
472	Z92837	elegans]	0.51	′123506	PROTEIN (HPS)	7.6	
	ĺ	Mouse mRNA for		1	1		
1		Rec A-like protein	l	1			
1		MmRad51, complete	1		(AB014607) KIAA0707 protein	4.5	
473	D13803	cds	0.51	3327228	[Homo sapiens]	4.5	
·			1	1	. manuago I sub-sia-i		
1					(AE001299) hypothetical	4.4	
474	X07187	Pea hsp21 mRNA	0.51	3328678	protein [Chlamydia trachomatis]	4.4	

1	Nearest	Neighbor (BlastN vs. G	enbank)	Nagaga Mai-k	h-/DL W	
SEO			CHOAHK/	ivearest iveign	abor (BlastX vs. Non-Redundant F	roteins)
Ð	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUI
		CCAAT/enhancer-				+
		binding protein				+
1		delta=transcription			1	
1		factor CRP3 homolog			1	1
1		[human, prostate			1	1
1 1		carcinoma cell line	1		(D90911) apolipoprotein N-	i
		LNCaP, Genomic,			acyltransferase [Synechocystis	i
475	S63168	1594 nt]	0.51	1653215	sp.]	1.2
		Xenopus laevis C2- HC type zinc finger			(AF067520) PITSLRE protein	
		protein X-MyT1	Ī		kinase beta SV2 isoform [Homo	Ī
476	U67078	mRNA, complete cds	0.51	3850320	sapiens)	0.17
477		Homo sapiens GT198 mRNA. complete			HYPOTHETICAL 100.6 KD TRP-ASP REPEATS CONTAINING PROTEIN C2C6.04C IN CHROMOSOME	
477	L38933	ORF	0.51	3219965	I	0.059
478	1	Lycopersicon esculentum polygalacturonase I	0.50	NOVE		
		S.cerevisiae	0.30	<none></none>	<none></none>	<none></none>
		chromosome XI reading frame ORF				
479		YKR079c	0.50	<none></none>	<none></none>	<none></none>
		Oncorhynchus keta	i	· · · · · · · · · · · · · · · · · · ·		~
480		GF-II gene	0.50	<none></none>	<none></none>	<none></none>
- 1		Homo Sapiens, RP58	T			
40.		DNA for complete	1			
481	AJ001388 1	nRNA	0.50	<none></none>	<none></none>	<none></none>

	Nearest	Neighbor (BlastN vs. (Senhank)	Nagara N	hha (Diany W	
		Treignoof (Blastit vs.)	Jenoank)	Nearest Neig	hbor (BlastX vs. Non-Redundant)	Proteins)
SEC	- 4	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Homo Sapiens, RP58				IT VALUE
		cDNA for complete		 		
481	AJ001388	mRNA '	0.50	<none></none>	<none></none>	NONE
		P.occultum 23S		410112	ZNONES	<none></none>
1	•	ribosomal RNA,		j		
482	M86626	partial cds.	0.50	<none></none>	<none></none>	<none></none>
					CHROMOSOME ASSEMBLY	CHOINES
1			1	1	PROTEIN XCAP-E African	
	j	Sambucus nigra lectin	i e		clawed frog >gi 563814	1 1
1	1	precursor mRNA,			(U13674) XCAP-E [Xenopus	1
483	U76523	complete cds	0.50	1722856	laevis	3.2
						1 32
		Mus musculus striatin			(M63730) BPAG2 [Homo	1 1
484	AF031663	mRNA, complete cds	0.50	179521	sapiens	3.2
						
1	}	Haemophilus	ŀ			1 1
	ł	influenzae Rd section	İ			1 1
1		44 of 163 of the			(Z92829) F10A3.15	1
485	U32729	complete genome	0.50	3875699	[Caenorhabditis elegans]	0.65
1		Dictyostelium			HYPOTHETICAL 28.3 KD	
		discoideum clone			PROTEIN IN GBD 5 REGION] [
1		9.10 Tdd-3 and RED	ı		(ORF4) >gi 2120954 pir I39562	ŀ
486	AF067198	repetitive elements,	0.50		ORF4 - Alcaligenes eutrophus	1 1
1-00	Ar06/198	partial sequence Human interleukin 4	0.50	2494740	>gi 695274 (L36817) ORF4	0.008
		(IL-4) gene, complete	İ			1
487	M23442	cds.	040	1101 m		1
107	1/123442	cus.	0.49	<none></none>	<none></none>	<none></none>
1		Caenorhabditis				
j i		elegans POU	ŀ		(15000100)	1
]	1	homeobox protein	I I		(AF098499) contains similarity	
		CEH-18 (ceh-18)			to Saccharomyces cerevisiae	l l
488		mRNA, complete cds.	0.47	3786409	MAFI protein (GB:U19492)	
		Lycopersicon		. 3780409	[Caenorhabditis elegans]	8.9
	1	esculentum	j			- 1
489		polygalacturonase 1	0.45	<none></none>	<none></none>	<none></none>
		Yersinia		NOTAL D	CNONES	SNONES
	[,	enterocolitica wbb	I]	1
490	Z18920	gene cluster	0.41	<none></none>	<none></none>	<none></none>
]	Human mRNA for			GTORES	CNONES
		KIAA0230 gene,	ł		(M64793) salivary proline-rich	į
491	D86983	partial cds	0.35	206712	protein [Rattus norvegicus]	4e-05
	Ī				pro (Atalias noi regicus)	-70-03
		lelianthus tuberosus	- 1			Ì
f	L L	ectin 2 mRNA,	ļ			i
492	AF064030 c	omplete cds	0.33	<none></none>	<none></none>	<none></none>

137 1	* Nearest	Neighbor (BlastN vs. (Genhank)	Nones Nois	LL (DI V N D	
SEC		Transpired (Blase 1 Vs. C	Jenoank,	ivearest iverg	hbor (BlastX vs. Non-Redundant)	Proteins)
ID	`	N DESCRIPTION			·	
==	TACCESSION	Vitreoscilla sp. outer	PVALUE	ACCESSION	DESCRIPTION	P VALUI
-		membrane protein				
}			1	l		
		homolog gene,		1		1
1	i .	complete cds; Trp				į.
1	1	repressor binding			J	
1		protein gene, partial		i	HYPOTHETICAL 24.5 KD	ı
493	A F0(7002	cds; and unknown	1		PROTEIN IN NADB-SRMB	1
493	AF067083	genes	0.33	401553	INTERGENIC REGION	8.3
1	1	Papio hamadryas	1			
1	į.	anubis gene encoding				İ
494	Y15520	fertilin alpha-II	0.20			
177	113320	Terriffin alpha-11	0.29	2408049	(Z99164) hypothetical protein	3.1
1					ARYL HYDROCARBON	
ł	ļ				RECEPTOR NUCLEAR	l
1	l				TRANSLOCATOR	1
1	ŀ	Alestes sp.			HOMOLOG (DARNT)	1 :
1	1	ependymin mRNA.	1		(TANGO PROTEIN)	1
495	U33475	partial cds		*****	transcription factor [Drosophila	!
-	033473	Mouse DNA for 8-	0.28	3913078	melanogaster]	1.4
ł	į	oxodGTPase.				
496	D88356	complete cds	0.22	ANONT:		
		Methanococcus	0.22	<none></none>	<none></none>	<none></none>
l		jannaschii section 145	Į			
1		of 150 of the	l		(TIS1222) = 10 (Green	1 1
497	U67603	complete genome	0.22	2209261	(U51222) p40 [Streptomyces	
				2209201	halstedii	8.3
]		Malurus cyaneus	- 1		(U29131) Mg-chelatase subunit	
498	U82386	microsatellite McyU2	0.22	992631	[Synechocystis sp.]	0.54
		S.cerevisiae			To vincende y sua Sp. J	0.56
		chromosome X	- 1			
	į	reading frame ORF	1		1	1
499		YJR125c	0.21	<none></none>	<none></none>	<none></none>
. [Dictyostelium			4.01.62	CAOMES
		discoideum AX2	1			
		protein tyrosine	į			1
ł		kinase gene, complete	1		1	ľ
500	U64830	ds.	0.21	<none></none>	<none></none>	<none></none>
ł					3.0.12	210112
- [Tuman prostate-	j			ı
		pecific antigen (PA)	1		(X97918) gene 12.1	1
501	M24543	tene, complete cds.	0.21		[Bacteriophage SPP1]	6.0

B. taurus mRNA for thrombospondin Salarus mRNA for thrombospondin Salarus microsatellite	16.5	Nearest	Neighbor (BlassN ve	Genhark)	Nanna Mar	hhar (BlassV vs. W. D. L	
D ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION P VALUE DESCRIPTION DESCRIPTION P VALUE DESCRIPTION	2129 Ann 3	.vemest	THEIRIDOI (DIASUV VS.	Jenbank)	Nearest Neig	nbor (Blasta vs. Non-Redundant P	roteins)
B. taurus mRNA for thrombosondin Soz X87618 (partial) 2162 bp 0.21 2146000 2146000 2249087 (partial) 2162 bp 0.21 2146000 22499087 (partial) 2162 bp 0.21 2146000 227 22830) orf [Homo sapiens] 2.7	`	ACCESSION	DESCRIPTION	P VALUE	ACCESSION		P VALUE
B. taurus mRNA for thrombospondin Solution Soluti		<u> </u>					T
502 X87618 (partial) 2162 bp 0.21 2146000 tuberculosis] 3.5						tuberculosis] >gi 1694863 gnl PID e283373 (Z83018) hypothetical protein	
Status	502	X87618		0.21	21.46000	1	
Human germline immunoglobulin lambda light chain gene 0.21 2119158 procollagen type V alpha 2 - mouse >gi]309181 2.7			B.taurus microsatellite				
Solition Solition	303	271391		0.21	1354453	(U52830) ort [Homo sapiens]	2.7
Note	504	V \$7000	immunoglobulin lambda light chain	201			
PROTEIN IN ABF2-CHL12 INTERGENIC REGION Sqi 1078003 pir S52835 hypothetical protein YMR075w yeast (Saccharomyces cerevisiae) sqi 763022 (Z48922) unknown (Saccharomyces cerevisiae) sqi 763022 (Z48922) unknown (Saccharomyces cerevisiae) sqi 763022 (Z48922) unknown (Saccharomyces cerevisiae) 2.0 UDP GLUCOSE:GLYCOPROTEIN GLUCOSYLTRANSFERASE PRECURSOR (DUGT) glucosyltransferase - fruit fly (Drosophila sp.) glucosyltransferase - fruit fly (Drosophila sp.) glucosyltransferase precursor (Drosophila melanogaster) 0.003 Rattus norvegicus nonmuscle myosin heavy chain-A heavy chain-A heavy chain-A mRNA, complete cds 0.21 3880111 Cenefinder 0.002 Cenefinder 0.002 Cenefinder 0.002 Cenefinder 0.002 Cenefinder Cenefinder 0.002 Cenefinder C	304	A37808	gene	0.21	2119158	mouse >gi 309181	2.7
mitotic			Yanana laasia			PROTEIN IN ABF2-CHL12 INTERGENIC REGION >gi 1078003 pir S52835 hypothetical protein YMR075w	
Mycobacterium fortuitum plasmid pJAZ38 replication protein Rep (rep) gene. complete cds 0.21 AF086476 AF086476 AF087006 Mycobacterium fortuitum plasmid pJAZ38 replication protein Rep (rep) gene. complete cds 0.21 2499087 DiDP- GLUCOSE:GLYCOPROTEIN GLUCOSYLTRANSFERASE PRECURSOR (DUGT) glucosyltransferase - fruit fly (Drosophila sp.) glucosyltransferase precursor [Drosophila melanogaster] 0.003 Rattus norvegicus nonmuscle myosin heavy chain-A amrNA, complete cds. 0.21 3880111 Genefinder (Z81130) predicted using Genefinder 0.002 LRR47 protein - fruit fly (Drosophila melanogaster) >gil415947 (X75760) LRR47 [Drosophila melanogaster] 1e-06 Homo sapiens full length insert cDNA 509 AF086476 AF087006 AF087006 Complete sequence 0.20 NONE> NONE>	505	U95098	mitotic phosphoprotein 44	0.21	2497139	cerevisiae) >gi 763022 (Z48952) unknown	7.0
Drosophila melanogaster 0.003 0.			fortuitum plasmid pJAZ38 replication			GDP-GLUCOSE:GLYCOPROTEIN GLUCOSYLTRANSFERASE PRECURSOR (DUGT) glucosyltransferase - fruit fly (Drosophila sp.)	2.0
Rattus norvegicus nonmuscle myosin heavy chain-A mRNA, complete cds. 0.21 3880111 Genefinder 0.002 Rabbit mRNA for aminopeptidase N (partial) 0.21 630864 [Drosophila melanogaster] 1e-06 Homo sapiens full length insert cDNA clone ZD88F12 0.20 <none> <none> <none> <none> <none> <none> <none> <none></none></none></none></none></none></none></none></none>	506			0.21	2499087		0.003
Rabbit mRNA for aminopeptidase N Silvarial Silva	507		Rattus norvegicus nonmuscle myosin heavy chain-A			(Z81130) predicted using	
Rabbit mRNA for aminopeptidase N Solution Solutio	-70/	031403	ilikina, complete cas.	0.21	3880111		0.002
Homo sapiens full length insert cDNA clone ZD88F12 0.20 NONE> NONE> NONE> S10 AF077006 complete sequence 0.20 NONE>	508	 	aminopeptidase N	0.21	62004	(Drosophila melanogaster) >gi 415947 (X75760) LRR47	
length insert cDNA				0.21	030864	[Drosophila melanogaster]	1e-06
plasmid pHPM186. 510 AF077006 complete sequence 0.20 <none> <none> <none></none></none></none>	509	AF086476	length insert cDNA clone ZD88F12	0.20	<none></none>	<none></none>	<none></none>
	510	1	plasmid pHPM186.	0.20	< NONF>	<none></none>	NONE
511 X75480 E.gunnii CAD gene. 0.20 <none> <none> <none></none></none></none>							

7.10	Nearest	Neighbor (BlastN vs. G	ienbank)	Nearest Neig	ghbor (BlastX vs. Non-Redundant F	
SEQ		1	I I	TACATCS! TACIS	gnbor (Blusta vs. Non-Redundant F	Toteins)
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU
		T.aestivum				·
1		mitochondrial nad7				
1		gene for NADH		Ì	1	i
1	[dehydrogenase		ł		1
512	X75036	subunit 7	0.20	<none></none>	<none></none>	<none></none>
		E.coli genomic DNA,			4.0.0.	THORE
ì		Kohara clone			<u>'</u>	i
513	D90875	#422(55.5-55.8 min.)	0.20	<none></none>	<none></none>	<none></none>
		Caenorhabditis		: <u>-</u>	4.0.1.2	CHONES
1	ł	elegans cosmid				Į
ĺ		F59B8, complete				1
1		sequence				1
		[Caenorhabditis			· ·	
514	Z68343	elegans]	0.20	<none></none>	<none></none>	<none></none>
l .		M.musculus V alpha				410110
515	X62486	11.1 gene 5'-region	0.20	<none></none>	<none></none>	<none></none>
]					PHOSPHORYLASE B	
		1	ł		KINASE ALPHA	1
			ŀ		REGULATORY CHAIN.	1
1 1		}	1		SKELETAL MUSCLE	
[]	i		ľ		ISOFORM	1
1 1			1		(PHOSPHORYLASE KINASE	
			1		ALPHA M SUBUNTT)	
		Caenorhabditis	- 1		>gi 2135923 pir I38111	
516	A F0 40 4 7 1	elegans cosmid			phosphorylase kinase (EC	
516	AF040651	W04H10	0.20	1170683	2.7.1.38) - human >gi 791043	7.4
		Pseudomonas	1	•	<u> </u>	
ı		fluorescens PHA				1
i		depolymerase (phaZ)	- 1		(A PO16024) DG2 (DI II	i
517		gene, complete cds.	0.20	3721862	(AB016024) Pfj2 [Plasmodium	
		Human mRNA for		3721802	falciparum] LAMININ ALPHA-1 CHAIN	1.9
ı		KIAA0194 gene,			PRECURSOR precursor -	
518	_	partial cds	0.20	126363	human	0.65
					(AC004908) similar to	0.65
- 1	ļ	c-scr=pp60c-src,	j		ribosomal protein L23a; similar	
ı		dr=src downstream	1		to P29316 (PID:g132848)	ł
519		egion	0.20	4159887	[Homo sapiens]	0.52
		vius musculus Balb/c				0.52
Į	Įŧ	orain-specific kinase	Í			j
	1.	Bsk) mRNA.			(M64793) salivary proline-rich	- 1
520	U07357	omplete cds.	0.20	206712	protein [Rattus norvegicus]	0.51

建	Negrace !	Neighbor (BlastN vs. G	enhank)	Negrare Nainh	bor (BlastX vs. Non-Redundant P	roteins\
	i vearest i	TEISHOOF (DISSUT VS. O	endank)	ivearest iveign	our (BlastA vs. Non-Redundant P	(Cieins)
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
						†
		Penicillium thomii				
1 1		internal transcribed				j
		spacer 1, 5.8S				1
		ribosomal RNA gene				
		and internal			AMINO-ACID	
1	·	transcribed spacer 2,		:	ACETYLTRANSFERASE	1
		complete sequence;			Pseudomonas aeruginosa	1
		and 28S ribosomal			>gi 151036 (M38358) N-	
		RNA gene, partial	_		acetylglutamate synthase	İ
521	AF034460	sequence	0.20	114136	[Pseudomonas aeruginosa]	0.39
				• •	ASSOCIATING FACTOR 1 (B	-
} i					CELL-SPECIFIC	i
) I	,				COACTIVATOR OBF-1) (OCT	Ì
1 1	i		ì		BINDING FACTOR I) (BOB-	
		1	0		I) (OCA-B) Bobl, B-cell-	ł
1 1			. 1		specific - mouse	
					>gi 1881818 bbs 179852	1
1 1			Ţ		mBob l=B-cell specific	i
		Xenopus laevis]		transcriptional coactivator line	İ
		mitotic	1		J558L, Peptide, 256 aa]	1
		phosphoprotein 44	[>gi 1353792 (U43788) Oct	
522	U95098	mRNA, partial cds	0.20	2842674	binding factor I [Mus musculus] (AL032643) similar to	0.073
			ľ		Uncharacterized protein family	
l	•		ŀ		UPF0034. Double-stranded	
•			j		RNA binding motif; cDNA EST	
1			ì		yk489b3.5 comes from this	}
ľ			ł		gene; cDNA EST yk439g7.5	1
i		S.lividans groEL2	J		comes from this gene	
523	X95971	gene	0.20	3925277	[Caenorhabditis elegans]	4c-19
		Ovis aries			(
		vasopressin Vl	ŀ	<i>i</i> .	_	
	Į,	receptor (VIR) gene,	1	•	-	
524	L41502	complete cds	0.19	<none></none>	<none></none>	<none></none>
		K.pneumoniae				
·	· ·	oxalacetate	1			
ľ		decarboxylase alpha	- 1			
		subunit gene.	1			
525		complete cds.	0.19	<none></none>	<none></none>	<none></none>
ļ	1	Helicobacter pylori,	- 1			
- 1		strain J99 section 12	1			İ
526	1	of 132 of the	0.10	NONE	aNO. m	
220	~EW1431 (complete genome	0.19	<none></none>	<none></none>	<none></none>

REES	Name	Mainhhan (Director Co		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
17147-142-14	1 3000	Neighbor (BlastN vs. G	enbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant I	roteins)	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
							
	·	Pedicularis verticillata chloroplast DNA, intergenic region between trnT(UGU)					
527	D88084	and trnL(UAA)5'exon	0.19	<none></none>	<none></none>	<none></none>	
528	U67599	Methanococcus jannaschii section 141 of 150 of the complete genome	0.19	· <none></none>	<none></none>	<none></none>	
						- CTOINE	
529	J05500	Human beta-spectrin (SPTB) mRNA, complete cds.	0.19	<none></none>	<none></none>	NONE	
1 22	303500	M.mycoides ftsY	0.19	CHONES	CNONES	<none></none>	
530	V10122	gene homologue and gene encoding					
530	Y10137	hypothetical protein	0.19	<none></none>	<none></none>	<none></none>	
531	AF027174	Arabidopsis thaliana cellulose synthase catalytic subunit (Ath-B) mRNA, complete cds	0.19	<none></none>	<none></none>	<none></none>	
532	·	Mouse thymic stromal cell mRNA for TLSF-beta, complete cds	0.19	<none></none>	<none></none>	<none></none>	
533	AJ012585	Tetrahymena thermophila macronuclear gene encoding ribosomal protein L3, exons 1-2	0.19	<none></none>	<none></none>	<none></none>	
524		Brassica napus 5- enolpyruvylshikimate- 3-phosphate synthase	0.10				
534	X51475	gene	0.19	<none></none>	<none></none>	<none></none>	
525	Į:	Sambucus nigra hevein-like protein					
535		mRNA, complete cds	0.19	<none></none>	<none></none>	<none></none>	
536	1	chromosome X reading frame ORF	0.19	. · <none></none>	ANONT:	-NONTE-	
	2-7027	*******	V.17	CAUIAES	<none></none>	<none></none>	

	Neares	Neighbor (BlastN vs. (Genbank)	Negrace Maia	hbor (BlastX vs. Non-Redundant l	Dana i a s
SEQ		1	- CHOWARY	TACTICS! TAGE	I Von-Kedundant I	roteins)
B	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE
		H.sapiens pilot				+
537	X63741	mRNA	0.19	<none></none>	<none></none>	<none></none>
1		O.latipes mRNA for				14.011
538	Y11255	annexin max4	0.19	<none></none>	<none></none>	<none></none>
539	L63537	Oncorhynchus mykiss (clone Jb-10) beta-2 microglobulin (B2m) mRNA. complete cds.		<none></none>	<none></none>	<none></none>
		N.tobacum T92 gene				1210112
540	X70903	for auxin-binding protein	0.19	<none></none>	<none></none>	<none></none>
		Caenorhabditis			·	
541	116:060	elegans cosmid				
341	U61958	C25A8 Macaca fascicularis	0.19	<none></none>	<none></none>	<none></none>
542	V.000.00	fertilin beta mRNA,				
342	U33959	complete cds	0.19	<none></none>	<none></none>	<none></none>
543	Z49835	H.sapiens mRNA for protein disulfide isomerase	. 0.19	2113940	(Z95556) hypothetical protein	
			0.17	2113940	Rv2507 PROBABLE E4 PROTEIN	9.4
544	AF035458	Spinacia oleracea heat shock 70 protein			papillomavirus (type 1) >gi 61015 (X62844) E4 gene product [Pygmy chimpanzee	
-344	AF033438	protein, complete cds Tetrahymena	0.19	267293	papillomavirus type 1]	9.4
545	U23441	thermophila B internal deletion sequence.	0.19	3877185	(Z66563) F46C3.2	
		Pneumocystis carinii		3677163	[Caenorhabditis elegans] (AF052502) DA26 homolog	9.3
546		major surface glycoprotein	0.19	3548901	[Epiphyas postvittana nucleopolyhedrovirus]	9.3
547		Rat ankyrin binding glycoprotein-1 related mRNA sequence.	0.19	3337352	(AC004481) putative chromatin	9.1
540		Methanococcus jannaschii section 102 of 150 of the			(Y13585) serotonin receptor 4	
548	U67560	complete genome	0.19	3183689	[Cavia porcellus]	8.7
549	r P	Mus musculus pacteria binding nacrophage receptor MARCO mRNA.	0.19		(AF089083) complement	
	210724	ompiete cus.	0.19	3659853	component ClqB like protein	7.1

4444						
		Neighbor (BlastN vs. C	Genbank)	Nearest Neigh	hbor (BlastX vs. Non-Redundant Pr	oteins)
SEQ	· •					
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE
			T		(U58751) C07G1.7 gene	- TIEGE
		T			product (Caenorhabditis	
550	X66467	C.albicans sec 18 gene	0.19	1326385	elegans	6.9
1					DIHYDROPYRIMIDINASE	- 0.7
1		Syngaster lepidus 16S		j	(DHPASE) dihydropyrimidinase	
	Ĭ	ribosomal RNA gene,	ĺ		- rat	
551	AF003487	partial sequence	0.19	3122039	>gi 1378019 gnl PID d1010479	6.9
1	Í					
1	į.	Rat calmodulin-			1.	
1		sensitive plasma	1			
1		membrane Ca2+-	ŀ	Ī	hypothetical protein - fruit fly	
1		transporting ATPase		1	(Drosophila melanogaster)	•
552	J05087	(PMCA3) mRNA.			>gi 296434 (X68408) ORF	
332	303087	complete cds. Homo sapiens	0.19	422462	[Drosophila melanogaster]	5.3
1	1	glutamate			PD 00 101 = 11 = 1	
ł	Į.	oxaloacetate			PROBABLE E4 PROTEIN	į.
553	AF080464	transaminase	0.10	2004024	>gi 790898 position 3286. 3288	j
1 335	74 000404	Human MEK kinase	0.19	3024834	is first start codon; putative	5.3
		3 mRNA, complete			(1179993)	
554	U78876	cds	0.19	1710445	(U78083) unknown [Emericella	
	3.3070	Vigna radiata mRNA	0.19	1710443	nidulans	5.3
		for proton		-*-	(AP000002) 256aa long	ŀ
i i		pyrophosphatase.	l		hypothetical protein	I
555	AB009077	complete cds	0.19	3256922	[Pyrococcus horikoshii]	5.1
					[2 yi de de da ilettresiat]	
1 1			1		(AF125463) contains similarity	í
		Xenopus laevis	i		to BTB (also known as BR-	i
		mitotic	i		C/Ttk) domains (Pfam:PF00651)	- 1
		phosphoprotein 44	Í		Score=62.8, E=7.6e-15, N=1)	1
556	U95098	mRNA, partial cds	0.19	4226159	[Caenorhabditis elegans]	4.1
1 1	İ		}			
			1		(AL031583) 1-	Į.
	1		ł	,	evidence=predicted by content;	
					1-method=genefinder;084; 1-	İ
l	j	1	1		method_score=47.46; 1-	j
	j		i		evidence_end; 2-	- 1
' , j	Į,	Escherichia coli K-12	1		evidence=predicted by match; 2-	i
1	1	MG1655 section 282	1		match_uccession=SWISS-	
- 1		of 400 of the	1	1	PROT:P23792; 2-	
557	1	complete genome	0.19		match_description=DISCONNE	
	12.000332	complete genome	0.19	3645960	CTED PROTEIN.; 2-matc	4.0

3	Nearest	Neighbor (BlastN vs. C	ienhank)	Negross Nain	hbar (Plane)	
SEC		Transfer (Diabat va. e	T T	Memest Meig	hbor (BlastX vs. Non-Redundant Pr	roteins)
B	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-	1					Ι
					(AL031383) 1- evidence=predicted by content; 1-method=genefinder;084; 1- method_score=47.46; 1- evidence_end; 2-	
558	AE000392	Escherichia coli K-12 MG1655 section 282 of 400 of the complete genome	0.19	3645960	evidence=predicted by match; 2- match_accession=SWISS- PROT:P23792; 2- match_description=DISCONNE CTED PROTEIN.; 2-matc	
559	L81774	Homo sapiens (subclone 3_d1 from P1 H25) DNA sequence	0.19	4001725	(AB015981) MnhA [Staphylococcus aureus]	3.0
560	AL021108	Drosophila melanogaster cosmid clone 137E7	0.19	4001688	(AB015718) protein kinase [Homo sapiens]	3.0
561	AB001510	Carabus leptoplesioides mitochondrial DNA for NADH dehydrogenase subunit 5, partial cds	0.19	3758855	(Z98551) MAL3P6.11 [Plasmodium falciparum] (AE001326) Amino Acid	2.4
562	AF069696	Egernia stokesii clone EST1 microsatellite	0.19	3328994	(Branched) Transport [Chlamvdia trachomatis]	2.4
563	X64144	F.pringlei ppcA1 gene for phosphoenolpyruvate carboxylase Human	0.19	3242974	(AF069555) G protein-coupled receptor p2y3 [Meleagris gallopavo]	2.3
564		immunodeficiency virus type l gag polyprotein (gag) gene. partial cds	0.19	2257710	(U73041) resolvase-like protein [Thiobacillus ferrooxidans]	2.3
		Danio rerio Noich nomologue 3 mRNA.			dependant epimerase/dehydratase family; cDNA EST EMBL:C10103 comes from this gene; cDNA EST EMBL:D66400 comes from this gene; cDNA EST EMBL:D70143 comes from this gene; cDNA EST EMBL:D70143 comes from this gene; cDNA EST EMBL:D70143 comes from this gene; cDNA EST yk493h11.3	2.3
565		complete cds	0.19		comes from	1.8

220	Nearest	Neighbor (BlastN vs. (2	1		
		THEIRIOOF (Blash Vs. (Jenoank)	Nearest Neig	hbor (BlastX vs. Non-Redundant Pr	oteins)
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
	 				masquerade precursor - fruit fly	
		R.norvegicus mRNA			(Drosophila melanogaster) >gi 665545 (U18130) masquerade [Drosophila melanogaster] >gi 1095942 prf 2110286A	
566	Y12502	for factor XIIIa	0.19	2133693	masquerade gene	1.8
·		BB1=malignant cell expression-enhanced gene/tumor progression-enhanced gene [human, UM-UC-9 bladder carcinoma cell line.			(U77783) N-methyl-D-aspartate	
567	S82470	mRNA, 1897 nt]	0.19	2444026	receptor 2D subunit precursor	
568	U97408	Caenorhabditis elegans cosmid F48A9	0.19	542433	[Homo sapiens] 225K protein - Babesia bovis (fragment)	1.8
569	U10470	Pseudomonas fluorescens PHA depolymerase (phaZ) gene, complete cds.	0.19	3721862	(AB016024) Pfj2 [Plasmodium falciparum]	1.7
570	M88160	Ovis aries MAF214 locus polymorphic dinucleotide repeat .	0.19	1293816	(U56963) T13A10.5 gene product [Caenorhabditis elegans]	1.4
571	AJI31336	mRNA for pollen allergen (Hol 1 2, group II) > :: emb AJ131339 LIT13 1339 Lolium italicum mRNA for pollen allergen (Lol i 2, group II) > allergen (Poa p 2, group II) > :: emb AJ131338 TAE1 31338 Triticum nestivum mRNA for pollen allergen (Tri a 2, group II)	0.19	3880447	(AL032675) predicted using Genefinder	0.82
	5	S.cerevisiae ARG8		3000-147	Generalidet	0.82
572	X84036	ind CDC33 genes	0.19	3882041	(AJ010405) hypothetical protein	0.62

	Nearest N	Neighbor (BlastN vs. G	enbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant Pr	oteins)
SEQ						
Ð	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Human WD protein			mucin - human >gi 501033	
		IR10 pre-mRNA.			(U14383) mucin [Homo	
573	U57058	partial cds	0.19	631302	sapiens	0.60
				•		
		Penicillium thomii				
		internal transcribed			1	
		spacer 1, 5.8S				
		ribosomal RNA gene				
		and internal			AMINO-ACID	
		transcribed spacer 2.			ACETYLTRANSFERASE	
		complete sequence;			Pseudomonas aeruginosa	
	•	and 28S ribosomal		•. •	>gi 151036 (M38358) N-	
		RNA gene, partial			acetylglutamate synthase	0.25
574	AF034460	sequence	0.19	114136	[Pseudomonas aeruginosa]	0.35
		Xenopus laevis			alpha-2-adrenergic receptor -	
		mitotic			human name 'ADRA2R' [Homo	1
	1105000	phosphoprotein 44	0.19	105270	sapiens	0.27
575	U95098	mRNA, partial cds Homo sapiens	0.19	103270	hypothetical protein 3 -	0.27
		genomic DNA, 21q			Pseudomonas sp. (DSM 6898)	
		region, clone:			plasmid pKB740 >gi 45867	
576	AG001475	125H6N2	0.19	94977	(X66604) ORF3	0.16
270	AG001473	123110112				
					TRANSCRIPTION	
		Mouse IgG receptor			INITIATION FACTOR TEILD	
		(beta-Fc-gamma-RII)			135 KD SUBUNIT (TAFII-135)	
		gene, exons 9 and 10,			(TAFII135) (TAFII-130) of	
		clones lambda-			RNA polymerase II transcription	
577	M63284	Fc(3.2.93).	0.19	3024681	factor TFIID [Homo sapiens]	0.088
		Pseudomonas				
		aeruginosa orotate		-		
		phophoribosyl				
1 1	1	transferase (pyrE).			·	
[]		catabolite repression control protein (crc)			1	
		and RNasePH (rph)			(AF055904) unknown	
578	U38241	genes, complete cds	0.19	3044086	[Myxococcus xanthus]	0.052
10/0	030241	genes, complete eds	0.17	20-7000	pistil extensin-like protein	
		*			(clone pMGI4) - common	
		Lontra longicaudis			tobacco (fragment) >gi 19927	
		transthyretin intron 1.	_		(Z14015) pistil extensin like	
579	AF039734	partial sequence	0.19	322759	protein [Nicotiana tabacum]	0.030
		Xenopus laevis XL-				
		INCENP (XL-			1	
		INCENP) mRNA.				
580	U95094	complete cds	0.19	2147194	collagen - Paralvinella grasslei	0.002

	Nearest	Neighbor (BlastN vs. C	enbank)	Nearest Neio	hbor (BlastX vs. Non-Redundant P	roteins)
SEC			1	1	1011-Redundant P	· ·
В	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-	+	Drosophila		 		
58.1	AB004232	melanogaster mRNA for DAD polypeptide, complete cds	0.19	2498765	PEROXISOMAL MEMBRANE PROTEIN PEX16 lipolytica)	1
		Gallus gallus alpha-			(U41272) T03G11.6 gene	0.002
		globin gene domain 5'		·	product [Caenorhabditis	
582	AF098919	region	0.19	1086863	elegans	4e-05
		Helicobacter pylori, strain J99 section 18			(AL022018) 1- evidence=predicted by content; 1-method=genefinder;084; 1- method_score=165.48; 1- evidence_end; 2- evidence=predicted by match; 2-	
1		of 132 of the			match_accession=AA264666; 2-	
583	AE001457	complete genome	0.19	2924552	match_description=LD08351.5p rime LD Drosophila melanoga	2- 05
		Plasmid RP4 traE	- 5.12	2724332	Time LD Drosophila melanoga	3e-05
584	L10329	gene. 3' end; traD gene. complete cds; traF gene. 5' end.	0.19	3878117	(Z49068) mitochondrial carrier protein	8e-07
585	AE001155	Borrelia burgdorferi (section 41 of 70) of the complete genome Orf virus E10R	0.19	861276	(U28739) similar to TPR domains in e.g. yeast STI1 protein [Caenorhabditis elegans]	2e-12
586	·U49979	homolog gene, partial cds, and DNA polymerase gene, complete cds Xenopus laevis	0.19	3850072	(AL033385) dna-directed rna polymerase iii subunit [Schizosaccharomyces pombe]	le-15
587		RanGTPase activating protein	0.19	995714	(X91258) pid:e198503 [Saccharomyces cerevisiae]	40.16
588	AF061854	Schizosaccharomyces pombe Clr4p (clr4) gene, complete cds	0.19	3242750	(AC005164) match to ESTs AA731149 (NID:g2140138), AA731908 (NID:g2752719), AA287837 (NID:g1933519), AA262811 (NID:g1898382), and AA825820 (NID:g2899132)	4e-16
589	ļ,	S.cerevisiae CHS2 gene encoding chitin synthase.	0.18	<none></none>	· <none></none>	<none></none>

	Nearest	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEC		Talgitoti (Biasirv vs. C	Jenounk)	Nearest Neig	nbor (BlastX vs. Non-Redundant)	Proteins)	
B	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE	
<u> </u>		Xenopus laevis XL-			•	 	
1		INCENP (XL-					
1		INCENP) mRNA,		1		1	
590	U95094	complete cds	0.18	<none></none>	<none></none>	<none></none>	
		Caenorhabditis				1 4,0,0	
1	1	elegans cosmid	1			1	
591	AF067610	F41A4	0.18	<none></none>	<none></none>	<none></none>	
	j	Homo sapiens	i			+	
ł		gonadotropin-					
l	1	releasing hormone				1	
1		precursor, second					
l .	1	form (GnRH-II) gene,	-	l	· [
592	AF036329	complete cds	0.18	<none></none>	<none></none>	<none></none>	
1		H.sapiens				1310112	
1		mitoxantrone-					
Į		resistance associated				1	
593	Z49216	mRNA	0.18	<none></none>	<none></none>	<none></none>	
l						11.01.12	
		Torulopsis glabrata			i	1	
ł		mitochondrial DNA]	
ĺ		for tRNA-Thr,-His			1	1	
		and -Glu upstream of					
594	X02167	cytochrome b gene	0.18	<none></none>	<none></none>	<none></none>	
		R.communis					
		(Carmencita) Scr1				1 1	
		mRNA for sucrose			·	[[
595		carrier	0.18	<none></none>	<none></none>	<none></none>	
		Homo sapiens	7				
		(subclone 2_c9 from	j	•		1 1	
ر م		P1 H56) DNA			55 KD ERYTHROCYTE	1 1	
596	L81692	sequence	81.0	1346575	MEMBRANE PROTEIN	8.4	
	l.	Mana-dia	T				
- 1	a	Nocardia	1			1	
ı		actamdurans pcbAB	1	•			
[and pcbC genes for	Į		1	 	
1		Ilpha-aminoadipyl-L-	ı			ı I	
ſ		ysteinyl-D-valine	1	•			
j		ynthetase and	1	•	SEED LIPOXYGENASE-2 (L-	ļ	
507		sopenicillin N			2) soybean >gi 170014 (J03211)	1	
597		ynthase	0.18	126404	lipoxygenase (EC 1.13.11.12)	6.5	
		Sus scrota	1				
- 1	1-	arathyroid receptor	1		 	1	
598	1,	PTH) mRNA.			(X04647) collagen alpha-2(IV)		
248	U18315 c	omplete cds	0.18	1022323	chain [Mus musculus]	3.8	

	Nearest	Neighbor (BlastN vs.	Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant Pr	
SEQ					, see to the redundant Fi	Totems)
Ð	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE
├	 			ļ		
	1		1	j	VERSICAN CORE PROTEIN	
]	ł			l	PRECURSOR	I
1	Į.	Plasmodium	1		PROTEOGLYCAN CORE	Į
	1	falciparum DNA ***		į.	PROTEIN 2) (GLIAL	Í
	1	SEQUENCING IN	ļ	j	HYALURONATE-BINDING	J
	i	PROGRESS ***	1		PROTEIN) (GHAP) >gi 608515	
		from contig 3-85.	1		(U16306) chondroitin sulfate	
599	AL010158	complete sequence	0.18	2506816	proteoglycan versican V0 splice-	1
		complete sequence	0.18	2300816	variant precursor peptide u0002b protein -	3.7
	i				Mycobacterium tuberculosis	
	1]	•	tuberculosis]	
	ł			I	>gi 1694863 gn1 PID e283373	
		Bos taurus mRNA for]]		(Z83018) hypothetical protein	
		thrombospondin 1,	i i		Rv2968c [Mycobacterium	
600	AB005287	complete cds	0.18	2146000	tuberculosis]	2.9
		Drosophila			(AL031371) hypothetical	2.9
		melanogaster cosmid			protein SC4G2.06	Į.
601	AL021108	clone 137E7	0.18	3483032	[Streptomyces coelicolor]	2.9
		Danio rerio Notch	j			ŀ
602	1153036	homologue 3 mRNA,			collagen alpha 1'(II) chain	l
002	U57975	complete cds	0.18	85719	precursor - African clawed frog	1.7
					(AL021387) similar to Zinc	
			1		finger, C4 type (two domains);	ì
ı			Í		cDNA EST yk452f4.5 comes	ı
ı		P.aeruginosa	1		from this gene; cDNA EST	
1	í	autonomously			EMBL:T00774 comes from this	ı
603		replicating sequence.	0.18	3878017	gene receptor NHR-3	
		repriedung sequence.	0.10	3676017	[Caenorhabditis elegans] STEM CELL PROTEIN	1.3
	}		į		chicken >gi 62845 (X63371)	
- 1					transforming capacity [Gallus	ł
604	X54965	G.sp alpha 5HR DNA	0.18	134304	gallus]	1.3
		Xenopus laevis			Sauras	1.3
	j.	mitotic	1		(X98893) hTAFII68 [Homo	i
		phosphoprotein 44	ſ		sapiens] splicing [Homo	- 1
605	U95098 I	nRNA, partial cds	0.18	1628403	sapiens]	1.3
	1.					
ŀ		Oryctolagus			92 KD TYPE IV	ł
		uniculus renal	1		COLLAGENASE	j
- 1		odium-dependent			PRECURSOR IV, 92K,	
1		hosphate transporter	!		precursor - rat >gi 1022784	
506		ype II mRNA.			(U36476) 92-kDa type IV	j
,50	020/93 [0	omplete cds.	0.18	1705984	collagenase (Rattus norvegicus)	1.2

	留 Neares	t Neighbor (BlastN vs. (Genhank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SE		1.5.Elloof (Dia301 V3. C	Jenoank)	Nearest Neig	nbor (BlastX vs. Non-Redundant P	roteins)
	`	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-⊢						
601	U23427	Human cholecystokinin type A receptor (CCK-A) gene, exons 1 and 2.	0.18	3261734	(Z94752) hypothetical protein Rv1004c	2.07
608	U49953	Rattus norvegicus protein kinase MUK2 mRNA, complete cds	0.18		(X81847) pectate lyase 1	0.97
609		Human alpha globin gene cluster on chromosome 16: zeta gene.	0.18	551238 1585259	[Erwinia carotovora] traJ gene [Amycolatopsis methanolica]	0.43
610	X62513	M.gallopavo gene for metallothionein	0.18	2494740	HYPOTHETICAL 28.3 KD PROTEIN IN GBD 5 REGION (ORF4) >gi 2120954 pir I39562 ORF4 - Alcaligenes eutrophus >gi 695274 (L36817) ORF4	0.31
611	X04862	Goat embryonic alpha globin gene zeta exons 2-3 Rat vitamin D	0.18	86837	androgen receptor B - human	0.082
612	M12450	binding protein (DBP) mRNA, complete cds.	0.18	4210432	(AJ130783) APC2 protein [Mus musculus]	0.038
613	AF038539	Mus musculus muscle NSP-like 1 (Nspl1) mRNA, complete cds	0.18	3297877	(AJ224868) GNASI [Homo	0.029
614		Bacteriophage P22 right operon, orf 48, replication genes 18 and 12, nin region genes, ninG phosphatase, late control gene 23, orf 60, complete cds, late control region, start of lysis gene 13	0.18		(U42436) C49H3.3 gene product (Caenorhabditis	
		of lysis gene 13 Pig mRNA for inter- alpha-trypsin nhibitor heavy-chain	0.18		(U61947) C06G3.8 gene	0.009
615	D38754 1	11. complete cds	0.18		legans	7e-06

	i Nearest	Neighbor (BlastN vs. (Tenbank)	Nagrace Maia	hhor (BlockY up No. B. 1	
SEC		TOTALION (DIESEL VS. C	Julionik)	ivearest iveig	hbor (BlastX vs. Non-Redundant Pr	oteins)
B	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
J					LRR47 protein - fruit fly	
1		Rabbit mRNA for			(Drosophila melanogaster)	
	l	aminopeptidase N	1	1	>gi 415947 (X75760) LRR47	ļ
616	X51508	(partial)	0.18	630864	[Drosophila melanogaster]	6e-07
1					HYPOTHETICAL IT.3 KD	
İ		-	1		PROTEIN C2C6.07 IN	
1	I				CHROMOSOME I	ĺ
1	ł	ĺ	1		>gi 2370504 gnl PID e339194]
1	i				pombe]	
1	1	S.kluyveri linear			>gi 3451305 gnl PID c1316730	
1	1	plasmid pSKL DNA	1		(AL031324) very hypothetical	
1	7,5,055	for open reading		٠.	protein [Schizosaccharomyces	
617	X54850	frames 1-10	81.0	3183405	pombe]	2e-08
l		i ·			(ALU32626) CDNA EST	
ı	İ				EMBL:D70654 comes from this	
1			i		gene; cDNA EST	·
	ł				EMBL:Z14359 comes from this	
1					gene: cDNA EST	
l	i				EMBL:D33409 comes from this	
1	1		j		gene; cDNA EST	
1	1	TT			EMBL:D36239 comes from this	- I
		Human peripheral	· [gene; cDNA EST	
618	L21954	benzodiazepine	0.10	300501	EMBL:Z14766 comes from this	ı
 	L21934	receptor gene, exon 4. Oryctolagus	0.18	3925211	gene	4e-09
		cuniculus protein	ŀ			ı
i		phosphatase 2A1 B	ľ			
1		gamma subunit			(11.07.1282)	
		(skeletal muscle			(AL034382) putative mitosis	f
		isolate) mRNA.	i		and maintenance of ploidy	i
619	U09355	complete cds.	0.18	3947877	protein [Schizosaccharomyces	
		Tomprete eas.		3541011	MYOSIN LIGHT CHAIN	8e-11
	1		i i		KINASE, SMOOTH MUSCLE	
•		T.cruzi hsp70 mRNA	j	-	AND NON-MUSCLE	j
		for 70 kDa heat shock	- 1		ISOZYMES (MLCK)	
620		protein, partial cds	0.18	3024081	(CONTAINS: TELOKIN)	9e-12
				3024001	(CONTAINS: TELORIN)	76-12
	i i	Mus musculus	1			
ı	[proteasome regulator			(U28739) similar to TPR	ì
	. 1	PA28 beta subunit			domains in e.g. yeast STI1	Ī
621		gene, complete cds	0.18	861276	protein [Caenorhabditis elegans]	le-14
		Methylobacterium			protein [Cachornabattis Cicgatis]	10-14
- 1		extorquens serine	j		(AF027208) AC133 antigen	1
622		cycle proteins	0.18	2688949	[Homo sapiens]	le-14
					[[.totalo sapietis]	10-14

	響 Neare:	st Neighbor (BlastN vs. C	Jenbank)	Nearest Neig	hbor (BlastX vs.: Non-Redundant F	
SE				THOU GOT THEIR	anoi (Diasox vs. Ivoli-Redundant F	roteins)
Щ	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
623	3 AF006573	Drosophila virilis maltase I (Mav1) and maltase 2 (Mav2) genes, complete cds	0.18	2500777	PUTATIVE RIBONUCLEASE III (RNASE III) >gi 3876420 gnl PID e1346063 (Z81070) similar to ribonuclease	
624		Staphylococcus aureus strain SA502A		2500558 <none></none>	[Caenorhabditis elegans]	2e-23
625	AJ223364			<none></none>	<none></none>	<none></none>
626	J03059	Human glucocerebrosidase (GCB) gene, complete cds	0.17	<none></none>	<none></none>	
627	AB008860	Fugu rubripes Ca12 gene for pheromone receptor, complete cds	0.17	2198849	(AF004900) E3KARP [Homo sapiens] >gi 2665826 (AF035771) Na+/H+ exchanger regulatory factor 2 [Homo sapiens] factor 2 [Homo sapiens] >gi 3618353 gnl PID d1034182 exchanger isoform A3 [Homo sapiens]	7.8
628	AF027174	Arabidopsis thaliana cellulose synthase catalytic subunit (Ath-B) mRNA, complete cds	0.17	539355	SCD25 protein (version 1) -	
629		Homo sapiens histone deacetylase 3 (HDAC3) gene, complete cds	0.17		hypothetical protein C15H7.1 - Caenorhabditis elegans	7.5 4.5

	源 Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant)	D : .
	Q D ACCESSIO		P VALUE			Proteins)
		DESCRIPTION	IFVALUE	ACCESSION	DESCRIPTION	P VALUE
	•			 	HYPOTHETICAL 83.2 KD	
-	1			}	PROTEIN F58A4.11 IN	
	Į.			1	CHROMOSOME III	1
		İ	1	1	>gi 3874287 gnl PID e1344088	
- 1	ł		1		EST EMBL:C12577 comes	
ı				i	from this gene; cDNA EST	
1	i	}	1	j	yk227e7.5 comes from this	1
	1		1	 	gene; cDNA EST yk303d1.5	1
	ì		1		comes from this gene; cDNA	
1	1		ļ		EST yk314c12.5 comes from	j j
	1			1.	this gene; cDNA	
1	ł		1 .]	EMBL:C11886 comes from this gene; cDNA EST	1
]	I	1		i	EMBL:C12577 comes from this	1 1
1			1		gene; cDNA EST yk227e7.5	}
	ľ	Homo sapiens histone			comes from this gene; cDNA	1 1
İ	i i	deacetylase 3			EST yk303d1.5 comes from this	1 1
630	AF059650	(HDAC3) gene,	1		gene; cDNA EST yk314c12.5	1 1
1000	AF039630	Chinese harnster	0.17	465932	comes from this gene; cDNA	4.4
		metallothionein II			(1.000.1.10)	
631	X55065	gene	0.17	3687237	(AC005169) putative Cys3His	
				3087237	zinc-finger protein cyclin E type II - fruit fly	1.5
	1	Rattus norvegicus			(Drosophila melanogaster)	
l	1 .	oxytocin receptor			>gi 429168 (X75027)	
632	1115290	(OTR) gene, exon 3	_		Drosophila cyclin E type II]
032	U15280	and complete cds Goat embryonic alpha	0.17	542565	[Drosophila melanogaster]	0.45
		globin gene zeta				
633	X04862	exons 2-3	0.17	86837		
		Plasmodium		80837	androgen receptor B - human	0.080
		falciparum DNA ***	í		(X95466) CPG2 protein [Rattus	1
 		SEQUENCING IN			norvegicus]	[
		PROGRESS ***	1		>gi 1588593 prf 2208498A	1
634	41.010222	from contig 4-09,			plasticity-related gene [Rattus	1
0.54	AL010222	complete sequence	0.17	1177322	norvegicus]	7c-07
		H.sapiens mRNA for	i			
635		MRP-1	0.17		(U92715) breast cancer	
		Orf virus E10R		3237300	antiestrogen resistance 3 protein	3e-09
ŀ		homolog gene, partial	1			j
ł	I	cds, and DNA	1	ĺ	(AL033385) dna-directed rna	i
476		polymerase gene,	1		polymerase iii subunit	- 1
636	U49979	complete cds	0.17		[Schizusaccharomyces pombe]	7e-15

WO 01/02568 PCT/US00/18374

	Nearest	Neighbor (BlastN vs. C	ienbank)	Nanzace Majak	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		T T T T T T T T T T T T T T T T T T T	T T	ivenest ivergi	ibor (Blasta vs. Non-Redundant Pr	oteins)		
В	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
	<u> </u>							
637	AF006573	Drosophila virilis maltase 1 (Mav1) and maltase 2 (Mav2) genes, complete cds	0.17	2500558	PUTATIVE RIBONUCLEASE III (RNASE III) >gi 3876420 gnl PID e1346063 (Z81070) similar to ribonuclease [Caenorhabditis elegans]	2c-29		
638	AE001141	Borrelia burgdorferi (section 27 of 70) of the complete genome	0.16	1850327	(U52370) fertilin beta [Homo sapiens]	2.3		
639	M72980	Anthonomus grandis vitellogenin gene (VTG), complete cds.	0.12	3242750	(AC005164) match to ESTs AA731149 (NID:g2140138), AA731908 (NID:g2752719), AA287837 (NID:g1933519), AA262811 (NID:g1898382), and AA825820 (NID:g2899132)	0.54		
640	AF023532	Simulium vittatum ATPase 6 gene, mitochondrial gene encoding mitochondrial protein, partial cds	0.11	<none></none>	<none></none>	2c-56		
641	U76523	Sambucus nigra lectin precursor mRNA. complete cds	0.10	3482965	(AL031369) putative protein	0.49		
642	AJ001596	Danio rerio mRNA for opioid receptor homologue	0.099	I7 06 694	LANOSTEROL SYNTHASE 5.4.99.7) - fission yeast (Schizosaccharomyces pombe)	2.3		
643	U26341	Oryctolagus cuniculus Na and Cl dependent betaine transporter mRNA, complete cds.	0.099	2645804	(AF033381) betaine homocysteine methyl transferase [Mus musculus]	0.59		
644	M11633	Bacteriophage Cp-5 (S.pneumoniae) 3' nverted terminal repeat. Streptomyces sp.	0.082		(AE000649) type IIS restriction enzyme R and M protein	4.3		
645	Į	gene for alkaline serine protease I	0.073		(U54641) 220 kDa silk protein [Chironomus thummi]	6.3		

	Nearest	Neighbor (BlastN vs. (Genhanki	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC		Treighbor (Brasus Vs. C	Conomik)	Nearest Neigh	noor (Blasta vs. Non-Redundant F	roteins)	
D	ACCESSION		PVALUE	ACCESSION	DESCRIPTION	P VALUE	
		Caenorhabditis					
646	Z72509	elegans cosmid F32G8. complete sequence [Caenorhabditis elegans]	0.072	<none></none>	NOVE		
1	2,230)	[CICEAID]	0.072	CHOINES	<none> (AC005954) ZO-3 [Homo</none>	<none></none>	
647	X70282	X.laevis xanf-1 gene	0.070	3851202	[AC003934) 20-3 [Homo [sapiens] [Homo sapiens]	0.40	
		Human DNA sequence from cosmid E141E2, on chromosome 22, complete sequence			Suprems (Promo suprems)	0.40	
648	Z69906	[Homo sapiens]	0.069	<none></none>	<none></none>	<none></none>	
649	AF056940	Drosophila virilis retrotransposon Tv1, complete sequence	0.069	2246532	(U93872) ORF 73, contains large complex repeat CR 73	5e-12	
650	AJ001151	Homo sapiens genomic sequence	0.068	<none></none>	<none></none>	<none></none>	
651	X54455	Bacteriophage BF23 gene 17 and gene 18	0.067	<none></none>	<none></none>	<none></none>	
652	X87936	P.pinea internal transcribed spacers 1 & 2 of ribosomal DNA	0.067	2459733	(U95374) aldehyde dehydrogenase [Haloferax volcanii]	4.3	
653		Dictyostelium discoideum TipD (tipD) gene, complete cds	0.067	3882275	(AB018320) KIAA0777 protein [Homo sapiens] IMETHIONINE	1.1	
654	X90592	O.cuniculus mRNA for p53 protein	. 0.067	1703275	AMINOPEPTIDASE 2 (METAP 2) GLYCOPROTEIN) (P67)	0.29	
655	U41805	Mus musculus putative T1/ST2 receptor binding protein precursor mRNA, partial cds Homo sapiens KIAA0421 mRNA,	0.067	642518	(U17326) neuronal nitric oxide synthuse [Homo sapiens]	0.29	
656		partial cds	0.066	<none></none>	<none></none>	NONE	
		Plasmodium Falciparum DNA *** SEQUENCING IN PROGRESS *** From contig 3-109.	5.535	CHOILES	<14014E>	<none></none>	
657	AL010213	omplete sequence	0.066	<none></none>	<none></none>	<none></none>	

1	Nearest	Neighbor (BlastN vs. C	i-nhank)	None	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SE		Thereinour Plaster vs. C	iciioank/	Nearest Neig	hbor (BlastX vs. Non-Redundant P	roteins)		
ID	~ j	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
						1		
		Homo sapiens mRNA				 		
658	AB014546	for KIAA0646 protein, complete cds	0.066	1082461	homeotic protein HB9 - human			
				1002.101	nomeone protein AB9 - numan	0.38		
	1	Rattus exulans isolate		i		1		
1	i .	huahine30 mitochondrial D-			(U24189) RRM-type RNA	1		
659	AF104156	loop, partial sequence	0.066	100000	binding protein [Caenorhabditis			
1000	74 104130	M.musculus mRNA	0.066	1002380	elegans]	0.29		
1	1	for spalt transcription		•	(47.005000)			
660	X97581	factor	0.066	. 4107313	(AL035075) putative myosin			
		Human clone H20 N-	0.000	410/313	heavy chain	0.28		
ł		acetylglucosaminyltra	٠.]		
.]		nsferase III DNA.			(U96963) p140mDia [Mus			
661	D85378	exon 2	0.066	2114473	musculus]	000		
					mascaras)	0.22		
	1	Human (clone	ı		{			
l		LA179) chromosome						
662	M97561	21 sequence.	0.065	<none></none>	<none></none>	<none></none>		
		Plasmodium						
		falciparum						
l		chromosome 2.	1			ĺ		
		section 10 of 73 of				j		
663	AE001373	the complete	0.05					
-003	AE001373	sequence growth hormone	0.065	<none></none>	<none></none>	<none></none>		
		receptor, growth	ļ	•	1	i		
		hormone binding	ł			1		
		protein {GHR/BP	1			1		
	1	gene) [mice, C57	- 1		1	i		
	l I	black/6, Genomic,	1		1	ŀ		
		179 nt, segment 8 of	- 1		7	j		
664		10]	0.065	<none></none>	<none;></none;>	<none></none>		
		Homo sapiens	 }-		210112	~10115		
	1	syntaxin 4 binding	İ		1	. [
1	, ,	protein UNC-18c	1	i	i	Į		
		UNC-18c) mRNA,	i		(AB006074) topoisomerase III	j		
665		complete cds	0.065	3061308	[Mus musculus]	0.82		
ı		vp[40]=svp-related	- 1					
ŀ		nuclear	1	İ	. }	- 1		
- 1	1	eceptor/retinoid	1		1			
		ignaling modulator zebrafishes, mRNA.	1		(U58734) weak similarity to	į		
666		876 nt]	0.065		ankyrin G (Caenorhabditis			
	200,00	0/0111	0.065	1326288	elegans)	0.12		

100		st Neighbor (BlastN vs.	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SE	- 1	DESCRIPTION	P VALUE	1			
			+	ACCESSION	DESCRIPTION HYPOTHETICAL 74.2 KD	P VALUI	
		G.domesticus mRNA for ventricular myosir			PROTEIN IN AMD1-RAD52 INTERGENIC REGION >gi 1077180 pir S49745 probable membrane protein YML034w - yeast (Saccharomyces cerevisiae) >gi 575685 (Z46659) unknown		
667	X59552	heavy chain	0.065	2497098	orf, len: 656, CAI: 0.13 [Saccharomyces cerevisiae]	0.014	
440	Vanaa	Anthonomus grandis vitellogenin gene			(AC005164) match to ESTs AA731149 (NID:g2140138), AA731908 (NID:g2752719), AA287837 (NID:g1933519), AA262811 (NID:g1898382),	0.014	
668	M72980	(VTG), complete cds.	0.065	3242750	and AA825820 (NID:g2899132)	5 e-33	
669	AB014546	Homo sapiens mRNA for KIAA0646 protein, complete cds	0.064	<none></none>	<none></none>	<none></none>	
670	M30039	Sheeppox virus strain KS-1 ORF HM1 gene, partial cds; ORF HM2 and ORF HM3 genes, complete cds; and ORF HM4					
0/0	W130039	gene, partial cds Caenorhabditis	0.064	<none></none>	<none></none>	<none></none>	
671	Z680i3	elegans cosmid W02H3, complete sequence [Caenorhabditis	anc:				
	200013	elegans]	0.064	<none></none>	<none></none>	<none></none>	
572	AF041332	Bodo saltans unknown mRNA, kinetoplast gene encoding kinetoplast protein. complete cds Mouse germline IgG-	0.064	<none></none>	<none></none>	<none></none>	
73	r	S chain gene, D-J-C egion, and switch egion.	0.064	<none></none>	<none></none>	<none></none>	

	Neares	Neighbor (BlastN vs. (Sambonk)	No.			
SEC		Treignbor (Diastry VS. (Jenoank)	Nearest Neig	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
B	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
<u> </u>	 						
674	U41289	Dictyostelium discoideum K7 kinesin-like protein mRNA, complete cds		3482972	(AL031369) putative protein	9,3	
675	M37395	L.lactis (strain SK11) proteinase plasmid pSK111 DNA, complete cds.	0.064	993019	(X87246) alternative start codon		
676	Z15030	H.sapiens gene for ventricular myosin light chain 2 > :: gb L01652 HUMVM LC Human ventricular myosin light chain 2 gene, seven exons.	0.064	730343	PROLACTIN RECEPTOR PRECURSOR (PRL-R) mouse >gi 220576 gnl PID d1001535 (D10214) prolactin receptor precursor [Mus musculus] >gi 293770 (L14811) prolactin receptor [Mus musculus] >gi 347842 (L13593) prolactin receptor [Mus musculus] receptor:ISOTYPE=long form [Mus musculus]	9.1	
677	Z12021	G.max gene for catalase Entamoeba histolytica	0.064	2498711	ORIGIN RECOGNITION COMPLEX PROTEIN, SUBUNIT 2 >gi 1185461 (U38472) essential ORC2- related fission replication factor Orp2 [Schizosaccharomyces pombe]	5.3	
678		protein serine/threonine kinase (pstk1) gene. complete cds.	0.064	733140	(U22453) carboxypeptidase [Simulium vittatum]	5.3	

4	Meare	st Neighbor (BlastN vs.	Genbank)	Naneage Main	the (PlaceV as No P.)	
SE				Treatest (4e)	phbor (BlastX vs. Non-Redundant Pr	oteins)
п	7	DESCRIPTION	P VALUE	4.00000000		
 	, icc255ic	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE
			- 			
	j	j			HYGROMYCIN-B KINASE]
	· I			1	(HYGROMYCIN B	i i
					PHOSPHOTRANSFERASE)	
		.	Í		(APH(7"))	1
	ł		1		>gi 66885 pir WGSMHH	
· [[1	hygromycin B	
	ł			1	phosphotransferase (EC 2.7.1)	
-	1			1	Streptomyces hygroscopicus	
1	ı				>gi 581682 (X03615) pot. hyg	1
1	1	* .		1 .	protein [Streptomyces	
1	1				hygroscopicus]	
	ı		٠.	1	phosphotransferase [synthetic	
	1	1	1	1	construct] >gi 2739064 cloning	
	1	Mus musculus alpha-	1		vector] >gi 2739068	i
		galactosidase A gene,			(AF025747) hygromycin B	ł
679	U50715	complete cds	0.064	125398	phosphotransferase [unidentified cloning vector]	
		S.cerevisiae	0.00	123338	Om(2D) protein - fruit fly	2.3
1	İ	chromosome XI			(Drosophila ananassae)	
	1_	reading frame ORF	İ		>gi 4-13770 gn1 PID d1006095	ļ
680	Z28182	YKL182w	0.064	1079035	(D26553) ORF	1.8
		Human ornithine	1 1			
681	M29917	aminotransferase	000		(U97553) unknown [murine	ł
1 30.	1/12/9/1/	gene, exon 1.	0.064	2317934	herpesvirus 68]	1.4
	Į.	Homo sapiens mRNA	1		(H20154) T07F12.2	
		for KIAA0902	}		(U29154) T07F12.3 gene product [Caenorhabditis	
682	AB020709	protein, complete cds	0.064	861404	elegans)	0.47
1						0.47
İ	1	Homo sapiens mRNA]	
407	1 200 1515	for KIAA0646			HOMEOBOX PROTEIN HB9	
683	AB014546	protein, complete cds	0.064	1708118	>gi 507425	0.35
1		Homo sapiens mRNA for NORI-1, complete	l l			
684	AB010427	icds	0.064	2200/7/	(AF015539) precollagen P	
			0.004	2388676	[Mytilus edulis]	0.018
		Orf virus ankyrin-like	[
		repeat protein, F11L			SSFI PROTEIN	j
	l l	homolog, and F12L	- 1		>gi 626624 pir S46700 SSF1	
		homolog genes.			protein - yeast (Saccharomyces	
685	U34774	complete cds.	0.064	731668	cerevisiae)	1e-05
		Mus musquiss	[
I		Mus musculus neuropilin-2(25)	i			i
ı		mRNA, alternatively			(AF073359) benzaldehyde	
686	AF022861	spliced, complete cds	0.064		dehydrogenuse [Pseudomonas	
		spireca, complete cus	0.004	4091978	sp. DJ77]	le-05

		Neares	t Neighbor (BlastN vs.	Genbank)	Noonet Vai	hhar (Diag V	
S	EQ			1	ivearest ivei	hbor (BlastX vs. Non-Redundant F	roteins)
	D)	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-						(AC004010) similar to Leucine	
6	87	U14331	Sus scrofa myogenin gene. complete cds	0.064	2781386	rich transmembrane proteins; 44% similarity to U42767 (PID:g1736918) [Homo	
			Chironomus pallidivittatus clone 1219 non-telomeric	0.000	2/81380	sapiens	3e-33
68	88	AF074870	Ssp repeat sequence	0.063	<none></none>	<none></none>	<none></none>
	39	705600	H.sapiens repeat			THORIE .	CHOINES
100	**	Z25523	region DNA. Plasmodium	0.063	<none></none>	<none></none>	<none></none>
-			falciparum chromosome 2, section 15 of 73 of the complete	٠.			
69	0	AE001378	sequence	0.063	<none></none>	<none></none>	<none></none>
69	1	Z 72947	S.cerevisiae chromosome VII reading frame ORF YGR 162w	0.063	<none></none>	<none></none>	<none></none>
692		Y14723	Choanomphalus incertus mitochondrial cytochrome c oxidase				ZIONES
"	+	114/23	subunit I gene, partial	0.063	<none></none>	<none></none>	<none></none>
603		2	Streptomyces sp. gene for alkaline	·		HYPOTHETICAL 108.5 KD PROTEIN IN UME3-PUB1 INTERGENIC REGION >gi 2131866 pir S62935 hypothetical protein YNL023c - yeast (Saccharomyces cerevisiae) >gi 1301855 gnl PID e239870 (Z71299) ORF YNL023c	
693	+-	X74103	serine protease I	0.063	1730713	[Saccharomyces cerevisiae]	6.7
			Iomo sapiens			GLUTATHIONE S- TRANSFERASE GST-6.0 (GST B1-1) >gi 421198 pir S29772 glutathione transferase (EC 2.5.1.18) - Proteus mirabilis >gi 2126142 pir S71882 glutathione transferase (EC	
			prouty 2 (SPRY2)	-		2.5.1.18) B - Proteus mirabilis	1
694	A	F039843 m	RNA. complete cds	0.063		egi 1053076 (U38482)	5.2

	Neares	Neighbor (BlastN vs. (Genbank)	Nearest Nein	hbor (BlastX vs. Non-Redundant Pr	
SEC		T		rearest rect	ibbi (Blastic Vs. Non-Redundant Pr	Oteins)
В	-	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-	 	<u> </u>	ļ			
					FRAGILE X MENTAL RETARDATION SYNDROME RELATED PROTEIN 2 >gi 2135129 pir S60173 fragile X mental retardation syndrome	
695	M63650	Mouse M-twist gene mRNA, complete cds.	0.063	1730141	related protein - human >gi 1098637 (U31501) fragile X mental retardation syndrome related protein [Homo sapiens]	1.0
696	V12200	Homo sapiens GDP dissociation inhibitor			hypothetical protein 4 - fowl	1.8
1 330	Y13298	beta pseudogene	0.063	1085930	adenovirus 1	1.3
697	X56600	Rat SOD-2 gene for manganese- containing superoxide dismutase	0.063	2000.40	(AB018254) KIAA0711 protein	
	7230000	disilidase	0.063	3882143	[Homo sapiens] HUNTINGTIN	0.60
698	Z 23107	M.musculus mRNA for 5HTx serotonin receptor	0.063	1700169	(HUNTINGTON'S DISEASE PROTEIN HOMOLOG) (HD	
	223107	Plasmodium vivax	0.063	1708162	PROTEIN)	0.45
699	M20670	circumsporozoite protein gene, 3' end.	0.063	4033395	DNA GYRASE SUBUNIT B subunit [Myxococcus xanthus]	0.35
700	• (H.sapiens CpG DNA, clone 76g11, reverse		10500	RETINOIC ACID RECEPTOR RXR-BETA sapiens] >gi 3172498 (AF065396) retinoic X receptor B dJ1033B10.11 (Retinoid X receptor beta (RXRB)) [Homo sapiens] >gi 4249766 (AF120161) retinoic X receptor	
-,00		read cpg76g11.rt1a . Xenopus laevis XL-	0.063	1350911	beta	0.16
701		INCENP (XL- INCENP) mRNA.	0.063	2981200	(AF048732) cyclin T2b [Homo	
	1	Xenopus laevis mitotic phosphoprotein 44	0.003	2901200	(Z81555) predicted using	0.090
702	1	nRNA, partial cds Yenopus laevis XL- NCENP (XL- NCENP) mRNA.	0.063	3877951	Genefinder (AL031174) hypothetical	6e-07
703		omplete cds	0.063	3393018	protein	2e-10

	※ Neares	t Neighbor (BlastN vs. (Genbank)	Nearest Neig	ghbor (BlastX vs. Non-Redundant)	
SE					groot (Blastic Vs. tvoll-Redundant)	roteins)
ID	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	
		E.coli genomic DNA.		1	Description	P VALU
1		Kohara clone			CYTOCHROME B561	
704	D90872	#419(54.7-55.1 min.)	0.063	2498198	(CYTOCHROME B-561)	3e-19
	Ī	M.crystallinum				1 30 17
		ferredoxin-NADP+		ł		
1	i	reductase (fnrA)				ŀ
705	M25528	mRNA, complete cds.	0.062	<none></none>	ANONT-	
				- CHOILE	<none></none>	<none></none>
		Strongyloides ratti	İ]
706	U45256	microsatellite B DNA	0.062	<none></none>	<none></none>	<none></none>
1	1	Xenopus laevis				
ł	ì	mitotic	,			
1		phosphoprotein 90	• •			1
707	U95102	mRNA, complete cds	0.062	<none></none>	<none></none>	NONT
					CNORES	<none></none>
Ī	1	Homo sapiens			<u>.</u>	
708	AF044317	TEL/AML1 fusion			İ	ĺ
1,00	71.044317	gene, partial sequence Caenorhabditis	0.062	<none></none>	<none></none>	<none></none>
İ		elegans cosmid				
l	1	T06E8, complete	ſ			İ
	1	sequence	1			
		[Caenorhabditis	1		(AC004663) Notch 3 [Homo	
709	Z73975	elegans]	0.062	3108187	sapiens]	2.9
		Human mRNA for			chitinase (EC 3.2.1.14)	
		heparan sulfate	İ		precursor - rice precursor - rice	
710		proteaglycan	0.062	1076741	>gi 807955 (X87109) chitinase	
			0.002	10/0/41	[Oryza sativa]	0.59
	ľ	Bovine retinal mRNA	1			i
 I		for transducin beta-		•	sialidase - Actinomyces viscosus	j
711		subunit	0.062	47.7578	>gi 141852	0.087
712		D.melanogaster Jun	0000			
		and 14-3-3 zeta gene Bombus terrestris	0.062	3879551	(Z70756) similar to collagen	0.073
l		nitochondrial			(U20600) NADH	
- [ytochrome oxidase I.			dehydrogenase subunit [Vanda	- 1
713		partial cds.	0.062	1684959	lamellata]	0.039

	Neares	t Neighbor (BlastN vs.	Genbank)	Nearest Maio	hhor (Plast V va Non D	
SEQ			1	Nearest Neig	hbor (BlastX vs. Non-Redundant)	Proteins)
B	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALL
<u> </u>					AMINOPERTIDASE B	—
			-		(ARGINYL	
		ł		1	AMINOPEPTIDASE)	
			1	Į.	(ARGININE	J
	i		1	1	AMINOPEPTIDASE)	ŧ
			1		(CYTOSOL	1
1			Į.	1	AMINOPEPTIDASE IV) (AP-	1
			1	1	B) >gi 2039143 (U61696)	1
714	175000	Human ladinin (LAD)	1	aminopeptidase B [Rattus	ŀ
/14	U58994	gene, complete cds	0.062	2811078	norvegicus]	9e-06
l			İ		(U58748) similar to potential	70.00
ł		Homo sapiens mRNA	.[transmembrane domains in S.	1
7,,	ADO: 455-	for KIAA0653	1]	cerevisiae nulcear division	
715	AB014553	protein, partial cds	0.062	1326350	RFT1 protein (SP:P38206)	5e-10
		j	1		HTPOTHETICAL 38.5 KD	1 33-10
I					PROTEIN IN ERVI-GLS2	1
1		1	1	J	INTERGENIC REGION	
1		1			>gi 2132587 pir S64322	ı
- 1					probable membrane protein	j
- 1					YGR031w - yeast	1
ı					(Saccharomyces cerevisiae)	1
		Mus musculus			>gi 1323010 gnl PID e243277	1.00
716	L16898	collagen alpha I type			(Z72816) ORF YGR031w	i
' ''	L10898	XVIII mRNA, 5'end. M.tuberculosis	0.062	1723657	[Saccharomyces cerevisiae]	le-14
717	77000.0	guaA/B & choD	0.040	_ \ /	(Z49907) B0491.1	
	2177343	genes	0.062	3873807	[Caenorhabditis elegans]	2e-19
		Homo sapiens MAD-				
		related gene SMAD7	J			
- 1		(SMAD7) mRNA.	j		1	
18		complete cds	0.061	A102 TO		
		Myrmeleon sp. 18S	0.061	<none></none>	<none></none>	<none></none>
19		ribosomal RNA.	0.061			
		10030Inat IQTA.	0.001	<none></none>	<none></none>	<none></none>
- 1	la	Choanomphalus	j	ł		
- 1		ncertus	- 1	j	1	
	- 1	nitochondrial	· 1		İ	
I		ytochrome e o xidase	1		· .	ľ
20		ubunit I gene. partial	0.061	-NIONTE		
		zene. puittut	0.001	<none></none>	<none></none>	<none></none>
	E	Bovine respiratory	Ì	1	· 1	
J		yncytial virus	ľ	1	1	1
		ucleoprotein mRNA.	ſ		ļ	i
21		omplete cds.	0.061	512055	1	
		- Pioto cus.	0.001	542955 r	nucleoporin p62 - human	8.6

SEC) I					TOTAINEL
ו ווי	· •				ghbor (BlastX vs. Non-Redundant P	(Clens)
<u> </u>	ACCESSIC	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	
					Sus scrota	P VALU
			T	 	>gi 494455 pdb TPOS B Sus	<u> </u>
1				l	scrofa Sus scrofa	1
				i	>gi 1421210 pdb 1PCP Porcine	1
	I		1	1	Spasmolytic Protein (Psp) (Nmr.	1
	1			Í	19 Structures) Spasmolytic	1
	ł	Xenopus laevis XL-	1	1	Polypeptide	l
	ł	INCENP (XL-	1	1 .	>gi 1633061 pdb 2PSP B Chain	
	I	INCENP) mRNA,	İ		B, Porcine Pancreatic	1
722	U95094	complete cds	0.061	494454	Spasmolytic Polypeptide	2.9
	i	Xenopus laevis			- Stypeptide	2.9
	•	mitotic			(AE001417) hypothetical	
723	1105000	phosphoprotein 44			protein [Plasmodium	
123	U95098	mRNA, partial cds	0.061	3845272	falciparum]	1.3
		Sporidiobolus salmonicolor	1 1			
		NADPH-dependent	1		ł.	
		aldehyde reductase	1 1		ĺ	
724	U26463	gene, complete cds	0.061		(U79302) unknown [Homo	
$\neg \vdash$	220,05	gene, complete cas	0.061	1710288	sapiens]	0.44
1					EMBL:D33048 comes from this	
					gene; cDNA EST	
- 1		ļ			EMBL:D35780 comes from this	
- 1	•	Į į	1		gene; cDNA EST yk442c6.3	
- 1			j		comes from this gene; cDNA	
- 1					EST yk442c6.5 comes from this	
- 1		[1		gene; cDNA EST yk398f6.3	
1			i		comes from this gene; cDNA	
- [1		E	
- 1			f		>gi 3979816 gnl PID e1358315	
ı			1		EST EMBL:D35780 comes	1
		1	1		from this gene; cDNA EST	l
- 1			.]		yk442c6.3 comes from this	Ī
		V	.]	•	gene; cDNA EST yk442c6.5	
- 1		Xenopus laevis weel homolog mRNA.	1		comes from this gene; cDNA	ı
25		complete cds	0.061		EST yk398f6.3 comes from this	
7		Caenorhabditis	0.061	3979720	gene; cDNA E	2e-04
I		elegans cosmid			HYPOTHETICAL 83.8 KD	
- 1		ZK1321, complete	1		PROTEIN C27F2.7 IN	ı
-		sequence	1		CHROMOSOME III	j
- 1		Caenorhabditis	1		>gi 1065510 (U40419) C27F2.7	l
6		elegans	0.061	3183491	gene product [Caenorhabditis	ſ

温暖		t Neighbor (BlastN vs. (Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant l	Proteins\
SEC					Enter (Diasex vs. 14011-Redundant)	roteins)
D	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	
	T		1	- RECESSION		P VALU
			-	 	HYPOTHETICAL 32.0 KD	
1	1	Zea mays pep gene			PROTEIN C09F5.2 IN CHROMOSOME III	1
1	l l	for (C3 type)	1	1	>gi 732538 (U22832) C09F5.2	i
1	j	phosphoenolpyruvate		J	gene product [Caenorhabditis	1
727	X61489	carboxylase	0.061	2496887	elegans]	1
l	1 .	Drosophila				le-15
]	1	melanogaster				1
728	A F005 400	Windbeutel (wind)	1		(AC005783) R33083_i [Homo	
/28	AF025408	gene, complete cds	0.061	3702295	sapiens]	2 c -60
1	İ	Brassica rapa mRNA	1			1
729	AB012106	for SRK45, complete				· ·
	AB012106	cds	0.060	<none></none>	<none></none>	<none></none>
		Arabidopsis thaliana				
	i	cellulose synthase			1	,
1	1	catalytic subunit (Ath-			·	
		B) mRNA, complete				l
730	AF027174	cds	0.060	<none></none>	NONE	
		H.sapiens mRNA for		4.02	<none> (AF077541) contains similarity</none>	<none></none>
		carnitine			to class-I aminoacyl-tRNA	
		palmitoyltransferase I			synthetases [Caenorhabditis	
731	Y08682	type I	0.060	3319446	elegans	8.1
ı		Xenopus laevis XL-	7			0.1
		INCENP (XL-	i			i
732	U95094	INCENP) mRNA.			(D78016) TRAE [Enterococcus	
	093094	complete cds	0.060	1041119	faecalis]	8.1
1		Helianthus tuberosus	1			
j		lectin 2 mRNA.	j		regulatory protein Rex - primate	ł
733		complete cds	0.060	632209	T-lymphotropic virus PTLV-L	
		Mus musculus	0.000	032209	(AF037401) neuropeptide	3.7
		Pontin52 mRNA,	J.		Y/peptide YY receptor Yc	
734	AF100694	complete cds	0.060	3098348	[Danio rerio]	1
- 1				50,00,0	LAR PROTEIN PRECURSOR	2.1
.			ŀ		(LEUKOCYTE ANTIGEN	
- 1	I_		1		RELATED)	
	P	Xenopus laevis			>gi 70146 pir TDHULK	
- 1		nitotic	- 1		leukocyte antigen-related	l l
735		phosphoprotein 90			protein precursor - human	j
, 33	U93102 In	nRNA, complete cds	0.060	125978	>2i 34267 sapiens]	1.2
l l		Sambucus nigra lectin	1			
		recursor mRNA.	1		(U87306) transmembrane	1
736		omplete cds	0.060		receptor UNC5H2 [Rattus	
		luman nuclear pore	0.060	2055394	norvegicus]	0.32
1		omplex-associated	1		(2) 10(2) (17)	
37		rotein TPR	0.060		(Y14063) ChTi thymocyte	
				7127034	antigen [Gallus gallus]	9c-04

	Negree	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC		INEIGHBOU (Blastin Vs. C	Jenoanki T	Nearest Neig	thor (BlastX vs. Non-Redundant P	roteins)		
B	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
 			I		(U58748) similar to potential			
- [1.	Homo sapiens mRNA			transmembrane domains in S.	1		
1	1	for KIAA0653		1	cerevisiae nulcear division			
738	AB014553	protein, partial cds	0.060	1326350	RFT1 protein (SP:P38206)	1e-09		
		Xenopus laevis	i					
1	1	mitotic			ļ.			
739	Linconn	phosphoprotein 44		1	(Y15513) Prodos protein			
139	U95098	mRNA, partial eds	0.060	2632098	[Drosophila melanogaster]	5e-10		
1		H.sapiens telomeric	İ	1	,			
	ļ	DNA sequence, clone			<u>'</u>	1		
ľ		12QTEL101, read				1. 1		
740	Z 96260	12QTELOO101.seq	0.059	<none></none>	21027			
		1100101011100	0.052	CNOINES	<none></none>	<none></none>		
1		Mouse homeobox						
1	i	protein (EVX2)				1 1		
741	M93128	mRNA, complete cds.	0.059	<none></none>	<none></none>	<none></none>		
		Brassica rapa mRNA			- CHOILE	CHOILES		
1		for SRK45, complete			(D90904) lysostaphin	1.		
742	AB012106	cds	0.059	1652318	[Synechocystis sp.]	4.7		
ĺ								
i .	1	Homo sapiens mRNA				i i		
743	4.7007000	for KIAA0451			transcription factor brn-3b -	}		
/43	AB007920	protein, complete cds	0.059	479491	human	0.71		
l .		Human histidine						
1		decarboxylase (HDC)	j		1			
744	M60445	mRNA, complete cds	0.058	NONE				
	17100143	mixiva, complete cus	0.038	<none></none>	<none></none>	<none></none>		
			ı		TRANSPORT SYSTEM			
l I			ł		PERMEASE PROTEIN OPPC]		
					>gi 1075086 pir D64184	ı		
	ſ		1		oligopeptide transport system	j		
			[•	permease protein (oppC)C	. 1		
	1	Ustilago maydis	1		homolog - Haemophilus	1		
İ		exodeoxyribonucleas	ł		influenzae (strain Rd KW20)	f		
		e (REC1) gene.	1		permease protein (oppC)	• 1		
745	U01836	complete cds.	0.058	1171908	[Haemophilus influenzae Rd]	1.5		
- 1	4,	L voonerrige -	1					
ļ		Lycopersicon	l l		j	l l		
ł		class II small heat			.	1		
		shock protein HCT2	İ		1	1		
		(HSP17.4) mRNA.	1	·	(15000131) 131	1		
746		complete cds	0.058	2103245	(AF069131) chitinase [Bacillus			
		Brassica rapa mRNA	0.000	3193265	subtilis	0.002		
i		or SLG45, complete	[(IIO3078) dunais bassus abais	j		
747		ds	0.057	433385	(U03978) dynein heavy chain isotype 7A [Tripneustes gratilla]	_ ,		
			5.057	40000	hisotype /A [1 inpineustes gratifia]]	3.4		

医		t Neighbor (BlastN vs. (Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant P	mtaise)
SE	-	N DESCRIPTION				ioteins)
F	TACCESSIO		PVALUE	ACCESSION	DESCRIPTION	P VALU
\vdash		MRNA for	 			
	j	neoxanthin cleavage	I			
748	AJ005813	enzyme	0.056			1
	12003013	Lagopus lagopus	0.056	<none></none>	<none></none>	<none:< td=""></none:<>
	1	genomic				
ł		microsatellite	1	[(A EQQUADO)	İ
749	Y16828	sequence, LLST4	0.056	3328678	(AE001299) hypothetical	[
			0.050	3328078	protein [Chlamydia trachomatis]	4.3
		Sambucus nigra		İ		1
1		ribosome inactivating		i	İ	1
İ	i	protein precursor				
750	AF012899	mRNA, complete cds	0.055	<none></none>	<none></none>	<none></none>
	1	,			69 KD PROTEIN	CHOINES
1		Sambucus nigra			>gi 94375 pir \$19150	1
751	45054005	hevein-like protein			hypothetical protein, 69K -	
131	AF074385	mRNA, complete cds	0.055	137339	turnip yellow mosaic virus	0.69
	l	Sambucus nigra lectin				
	1	precursor mRNA,				
752	U76523	complete cds	0.035	NONE		
	<u> </u>	complete cus	0.033	<none></none>	<none></none>	<none></none>
		Human retrovirus-like	İ			
753	M92069	sequence-isoleucine c	0.034	<none></none>	<none></none>	
		GIL=ankyrin-like		- Trongs	(AF043700) contains similarity	<none></none>
		repeat [orf virus OV.	(to human RNA-binding protein	
		NZ2, Genomic, 1608	í		FUS/TLS (SW:Q28009)	
754	S78516	nt]	0.033	2804465	[Caenorhabditis elegans]	0.15
					4-	0.13
1	!	Chialas	l		HYDROXYPHENYLPYRUVA	Į.
		Chicken myosin	- 1		TE DIOXYGENASE 4-	Í
ı		alkali light chain	- 1		hydroxyphenylpyruvate	ľ
755		mRNA, complete cds,	0.00-		dioxygenase [Mycosphaerella	ł
,,,,	14113040	lone pF1.	0.027	3334221	graminicola]	6e-17
- 1	L	Arabidopsis thaliana	·]			
- 1		ellulose synthase	1			1
- 1		atalytic subunit (Ath-]		1	ŀ
		B) mRNA, complete			(7060 10): 12: 1	1
756		ds	0.025		(Z96048) predicted using	
				3077013	Genefinder	5.0

	¥ Neares	t Neighbor (BlastN vs.	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC			J.	TYCLIEST TYCH	noor (BlastX vs. Non-Redundant I	roteins)	
B	-	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU	
—				ļ			
		Streptococcus		·	(268014) similar to ribose- phosphate pyrophosphokinase; cDNA EST EMBL:D73173 comes from this gene; cDNA EST EMBL:D70909 comes from this gene; cDNA EST EMBL:D73449 comes from this gene; cDNA EST		
757	AJ002291	pneumoniae pbp1b gene, complete	0.025	3880487	EMBL:D76167 comes from this	1	
758	X 79104	C.borulinum (NCTC 7272 type A) HA-33 and P-21 genes.	0.024	2648615	(AE000970) tungsten formylmethanofuran dehydrogenase, subunit B (fwdE 2) [Archaeoglobus fulgidus]	6.1	
759	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.024	1663698	(D83785) expressed ubiquitously; product similar to D.melanogaster mam protein. [Homo sapiens]	4.7	
760	U36197	Chlamydomonas reinhardtii cobalamin- independent methionine synthase mRNA, complete cds	0.024	585723	PEROXISOME PROLIFERATOR ACTIVATED RECEPTOR GAMMA (PPAR-GAMMA) >gi 283818 pir C42214 peroxisome proliferator- activated receptor gamma chain - African clawed frog >gi 214668 (M84163) peroxisome proliferator activated receptor gamma [Xenopus laevis]	0.42	
761		Macaca mulatta (clone MMVA63) T- cell receptor alpha (TCR A) mRNA. partial cds.	0.023	ANONIE.			
762	AF035948 1	Mus musculus insulin receptor substrate-3	0.023	<none></none>	SPLICEOSOME ASSOCIATED PROTEIN 49 spliceosome-associated protein SAP-49 - human >gi 556217	<none></none>	
763	ļ f	S.tuberosum mRNA for inorganic phosphate ransporter, StPT1	0.023	110072	proline-rich protein MP4 - mouse >gi 53182	0.18	

		st Neighbor (Blast:N vs.	Genbank)	Nearest Ne	ighbor (BlastX vs. Non-Redundant	Proteins)
SE	_``				, and a second and	roteins)
	ACCESSIO	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	DVALL
<u> </u>				T		P VALU
		L.esculentum mRNA				
76	4 X91212	for HD-ZIP protein	0.022	<none></none>	<none></none>	1
ı	1	Homo sapiens			KNONES	<none:< td=""></none:<>
1		chromosome 5, P1	1	i	i	ļ ·
1	1	clone 1209C1 (LBNI	-		1	į.
	ł	H104), complete	Į.	l	·	[
1	į.	sequence [Homo	1	1	1	1
76:	AC004498	sapiens]	0.022	<none></none>	1107=	
1		Human prostatic acid		410112	<none></none>	<none:< td=""></none:<>
	1	phosphatase (ACPP)	ł			
766	U07083	gene, exon I	0.022	· <none></none>		1
		S.tuberosum mRNA		KNONES	<none></none>	<none></none>
.1	I	for inorganic			1	
		phosphate				
767	X98890	transporter, StPT1	0.022	<none></none>		i
		L.esculentum LAT59	0.022	CHONES	<none></none>	<none></none>
l		gene 5'flanking	}			1
1	i	region, expressed				1
1	1	during pollen		•		1
768	X56488	maturation	0.022	-NONE-		1
		Pseudorabies virus	0.022	<none></none>	<none></none>	<none></none>
	1	with upstream and	1		1	
		downsteam				
769	M34651	sequences.	0.022	AMONIT		
		P.taeda gene for	0.0	<none></none>	<none></none>	<none></none>
		protochlorophyllide	i i		(70000 () 71	
7 70	X66727	reductase	0.022	2070517	(Z92806) K10G4.4	
		,	0.022	3878517	[Caenorhabditis elegans]	4.3
		Xenopus laevis	1		1	
		mitotic	1		100000	1 1
		phosphoprotein 90	1		(D89501) similar to salivary	; !
771	U95102	mRNA. complete cds	0.022	105.1450	proline-rich protein P-B [Homo	ļ 1
		Xenopus laevis	- 0.022	1854452	sapiens]	4.3
j		mitotic	1			
	I	Phosphoprotein 44	Į.		(A D005000) - (
772	****	mRNA, partial eds	0.022	3021699	(AB005298) BAI 2 [Homo	Ī
				3041099	sapiens]	0.64
1	Į:	H.sapiens XB gene	1		liver stage antigen LSA-1 -	
ı	I	for tenascin-X, intron	1		Plasmodium falciparum	1
773		14	0.022	627050	>gi 9916 (X56203) liver stage	1
\neg			<u> </u>	627059	antigen	0.058
[lo	Operfringens nunH	1		(1/3025)	1
774		ene & ORF1.2.3 & 4	0.022	2062407	(U78975) poly(ADP-ribose)	1
				-002407	glycohydrolase [Bos taurus]	0.056

	Neares	Neighbor (BlastN vs.	Genhank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC		Toolgander (25 table) vo.	T T	Treatest Iveign	nbor (Blasta, vs. Non-Redundant I	roteins)	
B	-	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
<u> </u>					UI SMALL NUCLEAR		
1	i				RIBONUCLEOPROTEIN 70	- 	
1				İ	KD (UI SNRNP 70 KD)	1	
1	ı		1		>gi 85864 pir S02016 U1		
į.					snRNP 70K protein - African	1	
ł	i		1		.clawed frog >gi 65179	1	
775	V: 4071	Galius gallus mRNA	l l		(X12430) U1 70K [Xenopus	1	
1 //3	Y14971	for K60 protein	0.022	134091	[laevis]	0.032	
1	1	Caenorhabditis	1		DNA REPAIR PROTEIN		
		elegans cosmid			RAD18 >gi 1150622 protein	i	
776	AF003133	T21E3	0.022	150000	rad18 [Schizosaccharomyces		
1.70	711 003133	121E3	0.022	1709997	pombe]	2e-08	
1	1	Caenorhabditis			DNA REPAIR PROTEIN		
1	ľ	elegans cosmid			RAD18 >gi 1150622 protein		
777	AF003133	T21E3	0.022	1709997	rad18 [Schizosaccharomyces pombe]	2.00	
		Human helix-loop-	0.022	1707777	pombej	2e-08	
1		helix proteins Id-1]			1	
1		(ID-1) and Id-1' (ID-	1			1	
1	1	1) genes, complete				1	
778	U57645	cds	0.021	<none></none>	<none></none>	<none></none>	
		Methanococcus				W.O.A.	
1	ł	jannaschii section 112	1 1		1		
1		of 150 of the	i I		1	1 .	
779	<u>U67570</u>	complete genome	0.021	<none></none>	<none></none>	<none></none>	
		Trypanosoma cruzi	j				
		calcium-binding	ļ			1 1	
780	L01584	protein (CUB2.8)	0.021	\		l I	
1,00	E01364	gene, complete cds.	0.021	<none></none>	<none></none>	<none></none>	
		Borrelia hermsii outer				j	
781	L04787	membrane lipoprotein	0.021	<none></none>	MONTE		
		Xenopus laevis XL-	<u> </u>	7,10,162	<none></none>	<none></none>	
i 1		INCENP (XL-	ł	•			
i i		INCENP) mRNA.	ĺ	į	·		
782		complete cds	0.021	<none></none>	<none></none>	<none></none>	
		Saashusa					
	1	Saccharomyces cerevisiae	1		i		
ı		mitochondrion	1	ļ		1	
1		ransfer RNA-Thri	1				
ĺ		tRNA-Thr) gene;	1	ł		I	
ł		ransfer RNA-Val	I.		i	1	
ŀ		tRNA-Val) gene;		Į	·	1	
1		oxi2 gene, complete	1	İ		1	
		ds; ORF2 and origin					
783		of replication (ori5).	0.021	NONE	· NONE		
		pireactor (OHD).	0.021	<none></none>	<none></none>	<none></none>	

建業	Neares	Neighbor (BlastN vs.	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SE	- 1	İ				1	
	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU	
-	+	Homo sapiens biliary					
1	1	glycoprotein (BGP)					
784	M76741	gene, partial cds.	0.021	MONE			
	1	Tetrahymena	0.021	<none></none>	<none></none>	<none></none>	
		thermophila histone		·			
1		H3 (HHT2) gene.		i	1.		
785	M87504	complete cds	0.021	<none></none>	<none></none>	<none></none>	
						CHONES	
		Human calpain-like	1 1]	ł	
706	******	protease (htra-3)			İ		
786	U94346	mRNA, complete cds	0.021	<none></none>	<none></none>	<none></none>	
[1	Trypanosoma cruzi					
	1	calcium-binding	· · ·				
787	101501	protein (CUB2.8)					
707	L01584	gene. complete cds. Pongo pygmaeus CT	0.021	<none></none>	<none></none>	<none></none>	
		microsatellite, clone	i 1				
	ł	#1, from the tandemly				1	
	i	repeated genes	1		1	ł	
		encoding U2 small					
		nuclear RNA (RNU2				1	
788	U36530	locus)	0.021	<none></none>	ANONT:		
	,	Human gene for		410112	<none></none>	<none></none>	
		interleukin 1 alpha			EARLY TRANSCRIPTION	1	
789	X03833	(IL-1 alpha)	0.021	416974	FACTOR 70 KD SUBUNIT	8.9	
						- 5.5	
	1	Dictyostelium]		ļ	1 1	
		discoideum guanine	Ţ		1	1 1	
		nucleotide-binding	Ī			! [
i		protein alpha subunit 5 (G alpha 5) gene,			(U58510) RNA helicase	l i	
790		complete cds.	0.001		homolog (Chlorarachnion	1 1	
	020800	complete cus.	0.021	1401211	CCMP621]	8.8	
- [-		1		ACONITATE HYDRATASE		
ı	la la	H.sapiens CpG DNA.	į		(CITRATE HYDRO-LYASE)	1	
		lone 13d2, reverse		•	(ACONITASE) >gi 2183256 (AF002133) aconitase		
791	1	ead epg13d2.rt1c.	120.0	3121732	[Mycobacterium avium]	7.0	
	I	Plasmodium		0121752	NINE PROTEIN	7.0	
ŀ		alciparum strain 7G8	· 		>gi 2120251 pir S66581		
- 1		hloroquine	1		hypothetical protein 56 - phage	- 1	
ſ		esistance candidate	1		82 >gi 1051114 (X92588)	İ	
l		rotein (cg2) gene.	1		orf56; related to nin60 (ninE) of	1	
792		omplete cds	0.021		bacteriophage lambda	5.8	
ł		lethanococcus	1				
[innaschii section 112	1		İ		
793		f 150 of the	0031		(AC000104) F19P19.17		
	00/3/0 (omplete genome	120.0	2341037	Arabidopsis thaliana]	4.0	

	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SI	EQ			Treatest Mel	Zilbor (BlastX vs. Non-Redundant F	roteins)	
	D ACCESSIO	ON DESCRIPTION	P VALUE	ACCESSION		P VALUE	
-					NUCLEAR FACTOR NF-		
	i	1			KAPPA-B P100 SUBUNIT	 	
	j				(H2TF1) (ONCOGENE LYT-	[
1	1		1	İ	10) (LYT10) [CONTAINS:	1	
75	Dosses	Human DNA for	1		NUCLEAR FACTOR NF-		
1-12	D86566	NOTCH4, partial cds	0.021	1708619	KAPPA-B P52 SUBUNIT	3.1	
	1	Streptomyces				1	
1	· ·	coelicolor sigma	1		hypothetical 119.5K protein	ł	
79	5 7 11640	factor (rpoX) gene,	1	1	(uvrA region) - Micrococcus	1	
1/3	5 L11648	complete cds.	0.021	· 79833	luteus	1.8	
					NEUROENDOCRINE	1	
	1	1	1		CONVERTASE I		
	1	V	1		PRECURSOR (NEC 1) (PC1)	}	
1	ı	Xenopus laevis XL-	j j		(PROHORMONE		
1	1	INCENP (XL-	1		CONVERTASE 1) propeptide	i •	
790	£ 1105004	INCENP) mRNA.	1 1		processing protease [Mus	. 1	
1 /3	6 U95094	complete cds	. 0.021	128000	cookii)	1.0	
	1	Rattus norvegicus	i l				
797	1120020	microtubule-			(X74416) beta-3 integrin		
1/9/	U30938	associated protein 2	0.021	468600	[Takifugu rubripes]	1.0	
1	1	Chicken mRNA for					
	1	TSC-22 variant,				j	
798	D82364	complete cds, clone	1		27 kda amelogenin		
1 1 7°	D02304	SLFEST52	0.021	693723	{alternatively spliced}	0.61	
	1	Caller - 11	1				
799	U40041	Gallus gallus eHAND		•	(AB011541) MEGF8 [Homo	į.	
	040041	mRNA, complete cds	0.021	3449308	sapiens]	0.21	
l	J	M saminas VD	l		liver stage antigen LSA-1 -		
l	J	H.sapiens XB gene for tenascin-X, intron	ł		Plasmodium falciparum	j	
800	X71932				>gi 9916 (X56203) liver stage	1	
 300	A/1932	Oryza sativa 24-	0.021	627059	antigen	0.054	
	1	methylene lophenol	ŀ	•			
	1	C24(1)methyltransfer				. 1	
	1	ase mRNA, complete	1	•		i	
801	AF042333	cds	0.001		(X83413) U88 [Human	- 1	
	74 042333	Las	0.021	854065	herpesvirus 6]	0.014	
	1	}	j				
	1	Rat apical endosomal	1	1		[
]	glycoprotein mRNA,	1	ł		J	
802	L37380		0.001		TRANSMEMBRANE	ł	
	637360	complete cds.	0.021		PROTEASE, SERINE 2	le-05	
	l j	Caenorhabditis	ľ		DNA REPAIR PROTEIN		
		· · · · · · · · · · · · · · · · · · ·	1		RAD18 >gi 1150622 protein	- 1	
		Alegans access J					
803		elegans cosmid T21E3	0.021		rad18 [Schizosaccharomyces		

Trace of	Nearest Neighbor (BlastN vs. Genbank)			Negreet Naighbur (D) and All Date II			
SEC	1.000	Treignoor (Diase 4 Vs. C	Jenbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
B	ACCESSIO		P VALUE	ACCESSION	DESCRIPTION	P VALUE	
	ļ	Rabbit mRNA for				1	
ı		calcium channel BI-2				 	
1	1	(lambda CBP109 and			(AL022117) hypothetical		
804	X57689	CB101)	0.021	2959370	protein	le-10	
					(U41534) coded for by C.	10-10	
		Xenopus laevis			elegans cDNA CEESI42F:	1	
1	Ĭ	mitotic	ŀ		Similar to helicases of	1	
1		phosphoprotein 90			SNF2/RAD54 family.	1	
805	U95102	mRNA, complete cds	0.021	1109830	[Caenorhabditis elegans] HYPOTHETICAL 38.5 KD	5e-11	
1							
1		'} .		ł .	PROTEIN IN ERVI-GLS2 .	1	
i	Ĭ			l "	INTERGENIC REGION	1	
1]		••	i	>gi 2132587 pir \$64322	1 1	
1	!				probable membrane protein	1 1	
1	[Į	;		YGR031w - yeast		
I			į		(Saccharomyces cerevisiae)		
					>gi 1323010 gnl PID e243277	1	
806	Vazzen	H.sapiens TROP-2			(Z72816) ORF YGR031w	ŀ	
1 800	X77753	S.tuberosum mRNA	0.021	1723657	[Saccharomyces cerevisiae]	5e-11	
1		for inorganic					
		phosphate				l i	
807	X98890	transporter, StPT1	0.021	2127070	zinc finger protein PZF - mouse		
	7,50050	transporter, Str 11	0.021	2137872	>gi 453376	2e-19	
1		Arabidopsis thaliana		•		} · ·	
1 1		cellulose synthase	1		!	i	
		catalytic subunit (Ath-	İ	·	· ·	1	
		A) mRNA, complete				i l	
808	AF027173	cds	0.020	<none></none>	<none></none>	ANONE	
		Homo sapiens	- 0.020	KITOITES	I KNOWES	<none></none>	
		Promotor Region and	ŀ			i	
809	AJ224935	PCK2 gene	0.020	<none></none>	<none></none>	<none></none>	
					3101122	2101123	
	j	Sambucus nigra	j		·		
		ribosome inactivating	- 1		ľ	1	
1		protein precursor	1	i		1	
810	U76524	mRNA, complete cds	0.020	<none></none>	<none></none>	<none></none>	
1		A.thaliana GBF1					
811	X99941	gene	0.020	<none></none>	<none></none>	<none></none>	
ł	1						
j	i i	M.musculus mRNA	ļ		ļ	J	
		for tyrosine kinase >	1			ł	
J		: gb S57168 S57168	•	1	1	l l	
]		Sek=Eph-related	1	l]	ł	
ı		eceptor protein	ŧ	-	J	1	
a, _ [yrosine kinase (mice.	i			l	
812	X65138 r	nRNA, 4242 nt]	0.020	<none></none>	<none></none>	<none></none>	

S.158:	Nearest Neighbor (BlastN vs. Genbank)			I No.			
		Neighbor (Blastin vs. C	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC	`	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
<u> </u>							
1		Borrelia hermsii outer	1				
813	L04787	membrane lipoprotein		<none></none>	<none></none>	<none></none>	
		Enterococcus faecium					
i		genes encoding					
	1	enterocin L50A and			1	l .	
ł	ł	enterocin L50B plus]	1	1	
814	A 10000 (00	5' and 3' flanking			}		
014	AJ223633	regions	0.020	<none></none>	<none></none>	<none></none>	
	1	Brassica rapa mRNA					
815	AB012106	for SRK45, complete	0.000	NONT	1	i . I	
1 813	ABUIZIUU	Helicobacter pylori,	0.020	<none></none>	<none></none>	<none></none>	
	1	strain J99 section 100	• • •		(T.11905)	1	
1		of 132 of the			(L11895) transmembrane	1	
816	AE001539	complete genome	0.020	172292	protein [Saccharomyces cerevisiae]		
1	112001337	complete genome	0.020	1/-472		8.4	
1	ŀ	Sambucus nigra			pol polyprotein - Chinese	i j	
		hevein-like protein	-		hamster intracisternal A-particle	1	
817	AF074386	mRNA, complete cds	0.020	94173	CHIAP34	8.0	
					C.I.I. U. 3-7	8.0	
	}	Herpesvirus saimiri					
	ł	dihydrofolate					
	•	reductase (DHFR)			Í		
ł		and snRNA (HSUR)			(Z98745) dJ29K1.2 [Homo		
818	M55264	genes, complete cds.	0.020	2924250	sapiens	6.5	
İ			- 1		D(4) DOPAMINE RECEPTOR		
1		· [I		(D(2C) DOPAMINE	1	
			1		RECEPTOR)	•	
	i		l		>gi 2119482 pir I49246 D4		
			- 1		dopamine receptor - mouse		
			ŀ		>gi 758427 (U19880) D4	İ	
					dopamine receptor [Mus		
		Homo sapiens clone			musculus]		
		24456 mRNA	1		>gi 1095539 prf 2109259A	l	
819		sequence	0.020	1706288	dopamine D4 receptor [Mus	40.	
	352103	Judgettee	0.020	1700200	musculus]	4.9	
		Sambucus nigra	1			į	
		hevein-like protein	ļ	Į	(Z83259) AmphiBrf38	I	
820		mRNA, complete cds	0.020		[Brunchiostoma floridae]	4.7	
					(Z67754) cDNA EST	- 4./	
			l l	1	EMBL:T02354 comes from this	- 1	
	ľ		ļ	ľ	gene; cDNA EST		
	1		- 1	1	EMBL:D32698 comes from this	- 1	
į		Homo sapiens clone	1		gene; cDNA EST		
į		24456 mRNA	ĺ	P.	EMBL:D35411 comes from this	- 1	
821	AF052163 s	equence	0.020	3874733	gene	4.7	

	Nearest	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ				remest recignor (Blascx vs. Non-Redundant Proteins)				
B	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU		
-		70 1 1 1 1				J		
1	1	Rat ankyrin binding						
822		glycoprotein-1 related			(K01664) Bkm-like protein	1		
822	L11002	mRNA sequence.	0.020	552132	[Drosophila melanogaster]	3.8		
1		Helicobacter pylori, strain J99 section 100		[}		
1	ľ	of 132 of the			(L11895) transmembrane			
823	AE001539	complete genome	0.020	172292	protein [Saccharomyces			
1000	142001333	complete genome	0.020	172292	(Correo) Summa to 11 K	3.8		
		!			Domain (2 domains); cDNA	1		
					EST yk390b10.3 comes from			
		l · ·			this gene; cDNA EST			
					EMBL:D71652 comes from this			
					gene; cDNA EST yk275f8.3			
		1	1		comes from this gene; cDNA	1		
İ				•	EST yk393b9.3 comes from this	ł		
]		1		l	gene; cDNA EST yk37	1		
					>gi 3880220 gni PID e1349842			
					yk390b10.3 comes from this			
			i		gene; cDNA EST	l		
			i		EMBL:D71652 comes from this	ļ		
		S.tuberosum mRNA			gene; cDNA EST yk275f8.3	1		
		for inorganic			comes from this gene; cDNA			
824	7/00000	phosphate			EST yk393b9.3 comes from this	٠.		
824	X98890	transporter, StPT1	0.020	3879798	gene; cDNA EST yk37	1.3		
					C-T-TETRAHYDROFOLATE SYNTHASE, CYTOPLASMIC			
		1			(C1-THF SYNTHASE)			
	f	1	- 1		(METHYLENETETRAHYDR			
ľ		1			OFOLATE			
- 1	Ĭ		1		DEHYDROGENASE /			
- 1	ł	Homo sapiens			METHENYLTETRAHYDROF			
ł		podocalyxin-like	1		OLATE CYCLOHYDROLASE			
		protein mRNA,			C1-tetrahydrofolate synthase	• .		
825	1	complete cds	0.020	1345633	[Rattus norvegicus]	0.066		
				<u></u>	DNA REPAIR PROTEIN	5.500		
- 1		Caenorhabditis	1		RAD18 >gi 1150622 protein			
		elegans cosmid	1	1	rad18 (Schizosaccharomyces			
826	AF003133	T21E3	0.020	1709997	pombe!	2e-07		
- 1	l,	T						
		Saccharomyces	1			1		
J	1	erevisiae VAR1	1					
Į	1-	gene, mitochondrial			1	ł		
ı		ene encoding	1			1		
		nitochondrial						
827		protein, 3' processing	2015					
02/	U32857 s	ite, partial sequence	0.019	<none></none>	<none></none>	<none></none>		

1 A	Neares	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC		T. S. Enoor (Diastia VS. C		ivearest Neis	gnoor (Blasta vs. Non-Redundant I	roteins)		
I ID		N DESCRIPTION	DVALUE	1				
—	TACCESSIO	N DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE		
-			ļ		NEUROGENIC LOCUS			
ı		A sahida asia shaliana	1	l .	NOTCH HOMOLOG			
1	1	Arabidopsis thaliana	i	Į	PROTEIN 4 PRECURSOR	Ţ		
ł	ł	cellulose synthase		1	(TRANSFORMING PROTEIN			
		catalytic subunit (Ath-	1	ł	INT-3) mammary gene mRNA,			
828	AF027174	B) mRNA, complete cds	0000		complete cds.], gene product			
1020	74.027174	cas	0.019	2506381	[Mus musculus] [ALU21481] similar to	3.3		
	I	1	l	ļ				
1	İ	Ì	•	ł	Phosphoglucomutase and phosphomannomutase	1		
1	1			·	phosphoserine; cDNA EST			
	i				EMBL:D36168 comes from this			
	Į.			i	gene; cDNA EST	· ·		
ł	i	Laccaria bicolor	٠.	l		1		
1		glyoxal malate			EMBL:D70697 comes from this	· [
ļ	J	synthase protein			gene; cDNA EST yk373h9.5	1		
829	AF034099	mRNA, complete cds	0.019	3880930	comes from this gene; cDNA EST EMBL:T008			
		Mus musculus	0.013	3000730	EST EMBL:1008	6e-15		
1	ł	Pontin52 mRNA.				1 1		
830	AF100694	complete cds	0.018	<none></none>	<none></none>	<none></none>		
		Human RP1 and			THOMES	ENONES		
	i .	complement C4B				1 1		
		precursor (C4B)			proline-rich protein precursor -			
831	U24578	genes, partial cds.	0.013	478673	kidney bean vulgaris]	3.1		
		C						
		Sambucus nigra lectin	ł		·			
832	U76523	precursor mRNA,	2011					
032	070323	complete eds	110.0	<none></none>	<none></none>	<none></none>		
		degrading bacterium	- 1		ĺ			
		DPO360 2.3-				1		
1		dihydroxybiphenyl						
		1,2-dioxygenase				İ		
		(bphC) gene.	. 1		[· .	. 1		
		complete cds and 2-	ŀ					
į		hydroxy-6-oxo-6-	l		1	į		
ŀ		phenylhexa-2,4-	ł			i		
		dienoic acid	1		·	i		
833	U57649	hydrolase	0.011	<none></none>	<none></none>	<none></none>		
I		Z.mays gene for		- 7		~1101112		
ł		phosphoenolpyruvate	1		. 1			
834	X15642	carboxylase	0.011	<none></none>	<none></none>	<none></none>		
1			T					
ا م		C.elegans collagen	I	i	(D83706) pyruvate carboxylase			
835		gene col-13	0.010	1695686	[Bacillus stearothermophilus]	3.1		
		Rattus norvegicus NF-	- 1					
836		KB gene. promotor			(AB020660) KIAA0853 protein	.]		
920	U83656	region	0.008	4240195	[Homo sapiens]	10.0		

PCT/US00/18374

- 20	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Neig	phoor (BlastX vs. Non-Redundant I	i
SE	Q				, serven Redundant	roteins)
Ш	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	
			i i	1	POL POLYPROTEIN	P VALU
1	1	Homo sapiens gene			(ORFIA/IB) (CONTAINS:	+
1		encoding retina-	1	ľ	RNA-DIRECTED RNA	· I
837	, , ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	specific guanylyl			POLYMERASE : HELICASE;	1
83	AJ222657	cyclase	0.008	417704	PROTEASE 1	7.4
1		1			CHLORIDE CHANNEL	
1	1			Į.	PROTEIN, SKELETAL	
	Ī		1	1	MUSCLE (CHLORIDE	1
	1		į	1	CHANNEL PROTEIN I) (CLC	-
1	ł	1	1		1) human >gi 397143 (Z25587)	
1	1				human CIC-1 muscle chloride	
1	1	Brassica rapa mRNA		• •	channel [Homo sapiens]	· ·
ł	ľ	for SRK45, complete			>gi 398161 (Z25884) human	1
838	AB012106		0.008	544024	CIC-1 muscle chloride channel	j
			0.000	344024	[Homo sapiens]	4.6
ł		Sambucus nigra			(U13643) similar to reverse	}
İ	I	ribosome inactivating			transcriptase; possible	
l		protein precursor			pseudogene [Caenorhabditis	i
839	U76524	mRNA, complete cds	0.008	532468	elegans]	3.8
						3.6
j	1	Sambucus nigra	Į.			
	1	ribosome inactivating			(AF002589) cytochrome	
840	AF012899	protein precursor			oxidase I [Austrofundulus	
070	AF012899	mRNA, complete cds	0.008	4101160	[imnaeus]	2.7
		Sambucus nigra	1			
		hevein-like protein				
841	AF074385	mRNA, complete cds	0.008	1211500	SRB-8/9 PROTEIN	
	3,000	ma va, complete cus	0.003	1711520	>gi 1334996	1.6
		Human non-muscle	1			
		alpha-actinin mRNA.	ı		(AC003301)	
842		complete cds	0.008	2829922	(AC002291) extensin	
		Homo sapiens 3-		2027322	[Arabidopsis thaliana]	0.11
į		hydroxyisobutyryl-			·	· · ·
- 1	ļ	coenzyme A				- 1
		hydrolase mRNA,				
843	U66669	complete cds	0.007	<none></none>	<none></none>	<none></none>
	l,	N/- DW: -	T			
		Mouse mRNA for	1			1
- 1		P100 serine protease	1	ļ		- 1
844		of Ra-reactive factor		Ì	ĺ	- 1
× 1	D10492 (RaRF), complete cds	0.007	<none></none>	<none></none>	NONE>

	. Ted Neare	Nearest Neighbor (BlastN vs. Genbank)			Negram Naighbor (DlassV N. D.			
	EQ	T Transfer (Brasil 173.	GCMGMIK)	Nearest Neigh	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	DACCESSIO	N DESCRIPTION	DYALLE					
F	- TACCESSIC	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE		
\vdash						T		
- 1	1	immunodeficiency		ł				
	Ī	virus type 1 proviral	į					
	1	DNA for envelope		1		1 1		
. 84	15 D90923	glycoprotein, partial cds. isolate 03S	0.007					
	270723	cus. Isolate 033	0.007	<none></none>	<none></none>	<none></none>		
	•	Homo sapiens mRN.A	.1					
	1	for KIAA0515	Ì			j i		
84	AB011087		0.007	<none></none>)	1		
			0.007		<none></none>	<none></none>		
		Aquifex aeolicus		• •		1		
	1	section 20 of 109 of	1			1 1		
84	7 AE000688		0.007	<none></none>	<none></none>	1 2/02/2		
		B.bovis WC1.1			CHOILES	<none></none>		
84	8 X63723	mRNA	0.007	<none></none>	. <none></none>	<none></none>		
						TATOM		
		Sambucus nigra		•	1	1		
	.=	hevein-like protein				1 1		
84	AF074386	mRNA, complete cds	0.007	<none></none>	<none></none>	<none></none>		
	1	Human beta globin				1 1		
850	J00097	region Alu repetitive	0.00-			1 1		
100	300097	sequence type T. Human	0.007	<none></none>	<none></none>	<none></none>		
1	1	immunodeficiency	1					
		virus type I proviral	1					
i i	1	DNA for envelope	1			1		
		glycoprotein, partial	ı			1 1		
851	D90923	cds, isolate 035	0.007	<none></none>	<none></none>	NOVE		
		Xenopus laevis XL-		41.014E2	ZNONES.	<none></none>		
l		INCENP (XL-	1			ľ		
	1	INCENP) mRNA,	l			i i		
852	U95094	complete cds	0.007	<none></none>	<none></none>	<none></none>		
		T.castaneum				31101112		
853	X91618	hunchback gene	0.007	<none></none>	<none></none>	<none></none>		
İ	ł	Rat nontranscribed						
	[spacer (NTS)			i	1		
854	V02020	downstream of 28S				1		
0.74	X03838	rRNA gene	0.007	<none></none>	<none></none>	<none></none>		
l	[Rattus pormaciona	ľ					
		Rattus norwegicus interleukin-2 receptor	1	İ		1		
		alpha chain (CD25)	1	}		i		
855		mRNA, complete cds.	0.007	NONE		1		
	1.230.45	marra complete cus.	0.007	<none></none>	<none></none>	<none></none>		

24	Neare	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SI	Q			Treatest Ivelg	- real est regiment (Blasck vs. Non-Redundant Proteins)			
	D ACCESSION	ON DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
-		Harris C. C. D.V.						
- 1		H.sapiens CpG DNA clone 9e2, reverse	٠	1				
85	6 Z64318	read cpg9e2.rt1a.	0.007			1		
	2,04318	read cpg9e2.ftfa.	0.007	<none></none>	<none></none>	<none></none>		
		Arabidopsis thaliana cellulose synthase catalytic subunit (Ath						
1	_	A) mRNA, complete	1			1 1		
85	7 AF027173	cds	0.007	<none></none>	<none></none>	<none></none>		
		Arabidopsis thaliana cellulose synthase catalytic subunit (Ath-B) mRNA, complete				CNONES		
858	AF027174	cds	0.007	<none></none>	<none></none>	<none></none>		
859	AF012899	Sambucus nigra ribosome inactivating protein precursor mRNA, complete cds	0.007		·			
1992	AI 012099	P.falciparum	0.007	<none></none>	<none></none>	<none></none>		
860	X95276	complete gene map of plastid-like DNA	0.007	<none></none>	<none></none>	<none></none>		
861	UZaas	Lycopersicon esculentum class II small heut shock protein Le-HSP17.6						
901	U72396	mRNA, complete cds	0.007	<none></none>	<none></none>	<none></none>		
862	AF100694	Mus musculus Pontin52 mRNA, complete cds	0.007	<none></none>	<none></none>	<none></none>		
		Leucania seperata nuclear polyhedrosis virus DNA for p13. xe, envelope protein.						
863	AB000383	complete cds	0.007	<none></none>	<none></none>	<none></none>		
864		Human DNA for NOTCH4, partial eds	0.007	<none></none>		<none></none>		
865		Sambucus nigra ribosome inactivating protein precursor mRNA, complete cds	0.007	VOVE				
		complete cus	0.007	<none></none>	<none></none>	<none></none>		

	Neares	Neighbor (BlastN vs. (Genhank)	Negros M	th - /D1 / N	
SE		1	T	IACTLES! IASIS	hbor (BlastX vs. Non-Redundant P	roteins)
II	- •	DESCRIPTION	P VALUE	ACCERTICAL	25002	
-	1	V DESCRIPTION	TEVALUE	ACCESSION	DESCRIPTION	P VALUE
		Arabidopsis thaliana	 			
- 1	ſ	cellulose synthase				
İ	ł	catalytic subunit (Ath	_	ł		1
- 1	ł	A) mRNA, complete	1		(AF058825) No definition line	
866	AF027173	cds	0.007	3047072	found [Arabidopsis thaliana]	1
				3047075	round (Arabidopsis manana)	8.9
i	1	Arabidopsis thaliana		ł		
1	1	cellulose synthase	1		1	!
	i	catalytic subunit (Ath-	-			
	I I	B) mRNA, complete			(U29359) SpaO [Salmonella	
867	AF027174	cds	0.007 .	975754	enterica]	8.6
1	j					- 5.5
	1	1	1		(U50199) coded for by C.	
	1	1			elegans cDNA yk89e9.5; coded	
1	i	j	[for by C. elegans cDNA cm7g5;	
}	İ	1			coded for by C. elegans cDNA	•
ł		Sambucus nigra			cm14b9; coded for by C.	
1		ribosome inactivating			elegans cDNA yk52g5.5; coded	i
i	<u> </u>	protein precursor			for by C. elegans cDNA	ď
868	U76524	mRNA, complete cds	0.007	1012552	yk76e5.5; coded for by C.	ı
		Matri, complete cas	0.007	1213557	elegans cDNA ykl3if11.5; c	8.4
			l		ISOASPARTATE(D-	- 1
1			j		ASPARTATE) O-	- 1
i			- 1		METHYLTRANSFERASE	
[1		(PROTEIN-BETA-	1
1					ASPARTATE	ľ
l	i i	j	ł		METHYLTRANSFERASE)	- 1
İ]	1			(PIMT) (PROTEIN L-	1
			1		ISOASPARTYL/D-	
		j	{		ASPARTYL	· [
]		i		METHYLTRANSFERASE)	
		Brassica rapa mRNA	1		methyltransferase [Drosophila	1
869		for SRK45, complete	0.00=		melanogaster] >gi 1171337	1
409		eds Rattus norvegicus	0.007	2499568	melanogaster]	8.3
		nomer- ic mRNA.	1		(A F005353) . 11 . 11	
870		complete cds	0.007	4002022	(AF095353) toll-like receptor 4	
		onipiete cus	5.007	4092077	mutant [Mus musculus]	6.2
ľ	S	Sambucus nigra				1
		evein-like protein	ł		(M80653) tetraheme	1
871		nRNA, complete cds	0.007	151377	[Pseudomonas stutzeri]	42
7			-		TRANSCRIPTION	6.2
		Bos taurus (clone	1		TERMINATION FACTOR	
872	L42319 S	al3.8) tristetraprolin	0.007		RHO	5.5
						٠.٠

11	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC	 	Treignoor (Diasis 1 43. C	I I	Mentezt Meiñ	noor (Blasta vs. Non-Redundant Pr	oteins)	
I ID	- I	DESCRIPTION				·	
<u> </u>	ACCESSION	1 DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
-	 	 	ļ				
ŀ	1	1	1	ł	(Z69637) Similarity to Human		
	Ĭ		1		Prolyi 4-hydroxylase alpha	l	
1					subunit (SW:P4HA_HUMAN);	ľ	
1					cDNA EST yk219g12.5 comes	1	
			ł		from this gene; cDNA EST		
	ļ	Human complement			yk319d8.5 comes from this		
I	į	component C4A	į		gene; cDNA EST yk339d11.5		
873	M59815	gene. exons 10	0.007	227.7.	comes from this gene; cDNA		
873	M139813	through 41. B.bovis WC1.1	0.007	3876769	EST yk371c9.3	5.3	
874	X63723	mRNA	0.007	7060000	(AJ001858) human SIM2		
F	A03/23	Expression vector	0.007	2969893	[Homo sapiens]	5.3	
ŀ		pME18S-FL3,			-45 NE E21- 15		
875	AB009864	complete sequence	0.007	2127610	p45 NF-E2 related factor 2 -		
19.5	AB007804	Complete sequence	0.007	2137618	mouse musculus]	5.1	
1	· ·	Sambucus nigra			1		
ļ		ribosome inactivating			1		
1		protein precursor			(AF043705) contains similarity		
876	U76524	mRNA, complete cds	0.007	2804497	to C2H2-type zinc fingers	5.0	
			5.007	2004477	to C2112-type zinc tingers	3.0	
l	ì	Xenopus laevis			i .		
		mitotic			(L27469) product of alternative		
		phosphoprotein 90			splicing [Drosophila	1	
877	U95102	mRNA, complete cds	0.007	440298	melanogaster]	4.7	
		Chicken mRNA for					
		aldehyde			(L75945) flagellar export	ĺ	
878	X58869	dehydrogenase	0.007	1185062	protein [Borrelia burgdorferi]	4.1	
		Nephila clavipes		<u> </u>			
		minor ampullate silk	i	•	(AF017434) pmi-like gene		
022	4 F00	protein MiSp1	1		product [Methylobacterium	ŀ	
879	AF027735	mRNA, partial eds	0.007	2394390	extorquens]	4.0	
- 1	ĺ						
880	AEIGEOGG	Bos taurus tuftelin					
000	AF105228	mRNA. complete cds	0.007	~ 3036802	(AL022373) putative protein HYPOTHETICAL 60.2 KD	3.9	
	ĺ		1		PROTEIN T27F2.1 IN		
	ļ	j	į		CHROMOSOME V	ì	
ı	1		ļ		>gi 3880311 gnl PID e1349855		
ı	1		- 1		BX42 (SW:BX42_DROME);	ļ	
1	į		1		cDNA EST EMBL:C07233	[
j			1		comes from this gene; cDNA		
l	}		1		EST EMBL: C08532 comes	1	
j	l,	Mus musculus]		from this gene; cDNA EST	J	
ł		Pontin52 ntRNA.			yk501h10.3 comes from this		
881	ı	complete cds	0.007	2500814	gene; cDNA EST yk501f1.3	20	
			V.007	-200014	gene, CDINA EST YKDUITT.S	3.8	

	Neare:	st Neighbor (BlastN vs. (Genbank)	Neurast Maio	hhar (Diany)	
SE				TACTICS! [AGIS	hbor (BlastX vs. Non-Redundant P	roteins)
П	ACCESSIO	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION:	ł
			1	ACCESSION	DESCRIPTION	P VALUE
		L.major mRNA for	 	 	(U78289) tylactone synthase	
88	2 X93567	beta-tubulin (1404bp.	0.007	2317862	modules 4 & 5 [Streptomyces fradiae]	
						3.0
	ł		ļ	ł	(AL032646) predicted using	
	İ				Genefinder; cDNA EST	
	İ		1	ł	EMBL:D76407 comes from this	
	1	·	1		gene; cDNA EST	
1	1		İ		EMBL:C08999 comes from this	
1	1	Brassica rapa mRNA		ł	gene; cDNA EST yk199b12.5	
		for SRK45, complete		٠.	comes from this gene; cDNA	
883	AB012106		0.007.	3881103	EST yk282a4.5 comes from this gene; cDNA EST EMBL:C0	
	1			3001103	THROMBIN RECEPTOR	2.7
	ı	Homo sapiens			PRECURSOR human	- 1
004	15041056	WSCR4 gene, exons			>gi 339677 (M62424) thrombin	.
884	AF041056	3 and 4	0.007	135817	receptor [Homo sapiens]	2.2
1		Rattus norvegicus			HYPOTHETICAL 32.2 KD	
1		homer-1c mRNA.	- 1		PROTEIN C22E12.04 IN	
885	AF093268	complete cds	0.007	1707510	CHROMOSOME I >gi 1220279	Ī
		Hevea brasiliensis 3-	0.007	1723518	(Z70043) unknown	2.1
ł		hydroxy-3-	ı			- 1
İ		methylglutaryl-	i		i [
	Į.	coenzyme A	1			
886	M74798	reductase gene. 3'	- 1			
000	W174798	end.	0.007	1001282	(D64003) polyA polymerase	1.9
1	l		1		HYPOTHETICAL III.9 KD	
		•	1		PROTEIN C34E10.8 IN	
		H.sapiens CpG DNA.	1		CHROMOSOME III	i
!		clone 76g11, reverse			>gi 500731 (U10402) weakly similar to protein C kinase	- 1
887	Z62997	read cpg76g11.rt1a.	0.007	1176532	substrate [Caenorhabditis	, . I
					DVA-I POLYPROTEIN	1.8
			ŀ		PRECURSOR nematode	1
	ŀ		1		polyprotein antigen precursor	i
		Sambucus nigra	i i	l	[Dictyocaulus viviparus]	[
	l i	hevein-like protein	1		>gi 1585421 prf 2124414A	
888		mRNA. complete cds	0.007		polyprotein antigen/allergen	- 1
		Synechocystis species	0.007	2498317	[Dictyocaulus viviparus]	1.2
I		(strain PCC 6803)	i	1		i
		irg.A gene. complete	1],	(AB018320) KIAA0777 protein	
889	L29426	ds.	0.007		[Homo sapiens]	1.1
					onio suprensi	1.1

19	Nearest	Nearest Neighbor (BlastN vs. Genbank)			hha (Planty)	
SEC				Nearest Neighbor (BlastX vs. Non-Redundant Protein		
I ID	- 1	DESCRIPTION	P VALUE	4.6672222		
	TACCESSIO.	T DESCRIPTION	FVALUE	ACCESSION	DESCRIPTION	P VALUE
-		Mus musculus DNA	 			
1		for prostaglandin D2	- 0	ł		
1	I	synthase, complete	1			1
890	D83329	cds	0.007	100.7		ł
1	1 203327	100	0.007	1001741	(D64004) hypothetical protein	0.97
	1				HYPOTHETICAL 11.6 KD PROTEIN IN NUT1-ARO2	ł
1	ľ	Brassica rapa mRNA			INTERGENIC REGION	ł
1 -		for SRK45, complete			PRECURSOR YGL149w -	
891	AB012106	cds	0.007	1723928	yeast (Saccharomyces	00.
			-	1,23,20	GLUTENIN, HIGH	0.94
	1		[MOLECULAR WEIGHT	
1	1	Sambucus nigra	l i		SUBUNIT 12 PRECURSOR	
		ribosome inactivating	• •		>gi 82606 pir A24266 glutenin	
1		protein precursor			high molecular weight chain 12	
892	U76524	mRNA, complete cds	0 .007	121452	precursor - wheat >gi 21779	0.79
1	i					9.75
1		Arabidopsis thaliana			ļ	1
1	ł	cellulose synthase				1 1
1	1	catalytic subunit (Ath-	I		1]]
893	AF027173	A) mRNA, complete cds	0.007		(U30294) ORF2 [Prevotella	1
1000	AI 02/1/3	cas	0.007	927287	ruminicola]	0.35
I			1		(U40061) contains similarity to	ı
1		H.sapiens IMAGE	1		transmembrane domains like	
894	Y11918	cDNA clone 26881	0.007	1055188	those found in sugar transporter	
		Mus Musculus	0.007	1033100	proteins	0.26
		alphaA-crystallin-			(AF083061) ABC transporter	1
895	L36827	binding protein [0.007	4063019	TliF [Pseudomonas fluorescens]	0.21
		Mus Musculus			- 1. J. Seadonionas muorescens	0.21
		alphaA-crystallin-	ļ		(AF083061) ABC transporter	ĺ
896	L36827	binding protein I	0.007	4063019	TliF [Pseudomonas fluorescens]	0.20
	1		T		1323.236101	
		H.sapiens CpG DNA.	}	•	.	
897		lone 54c10, reverse	1		1	1
89/	Z65719 I	ead cpg54c10.rtlu.	0 .007	1097307	HIC-1 gene [Homo sapiens]	0.20
					UTROPHIN (DYSTROPHIN-	
1	Ι,	Helianthus tuberosus			RELATED PROTEIN I)	i
1	1	ectin 1 mRNA,	- 1		(DRPI) (DRP)	j
898		omplete cds	0.007	117.6	>gi 284488 pir S28381 utrophin	1
		Aus musculus	0.007	1174915	protein) [Homo sapiens]	0.002
		athepsin S (CatS)	- 1		(1179721) DNIA E-11	
899		ene. exon 6	0.007	1707017	(U78721) RNA helicase isolog	, I
				1707017	[Arabidopsis thaliana]	0.001

- 4	Nearest Neighbor (BlastN vs. Genbank)			Negress Neighbor (Planty on New Park 1 2			
SEC	1.00.00	Z. Z. CO. I DIBSEL VS. I	Jenoank/	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
ID ID	•	N DECCRETATION			1		
۳	ACCESSIO		PVALUE	ACCESSION	DESCRIPTION	P VALUE	
-		Oryctolagus				T	
	ŀ	cuniculus	1				
		gp42/basigin/OX-		ļ	1	1	
900	1162200	47/HT7 mRNA,	1			l	
1900	U62398	complete cds.	0.007	2370494	(Z98944) hypothetical protein	2e-04	
		M.musculus glutathione reductase	ł	Į.		1	
901	X76341	mRNA.	0.007	25.0000	(AC005594) R26984_1 [Homo		
-~•	77/0341	Rat (lambda 20B0.5)	0.007	3513303	sapiens]	8e-07	
1	İ	M-type 6-				i	
	İ	phosphofructo-2-			· ·	1	
	1	kinase/fructose-2, 6-	ĺ			1	
902	M26215	bisphosphatase	0.007	3036809	(AI 022272)		
	1		0.007	3030809	(AL022373) putative protein (AB007902) HH0712 cDNA	6e-15	
	1			ĺ	clone for KIAA0442 has a 574-		
	1	Homo sapiens			bp insertion at position 1474 of		
	i i	KIAA0442 mRNA.	. 1		the sequence of KIAA0442.		
903	AB007902	partial eds	0.007	2662165	[Homo sapiens]		
			0.007	1002103	[[riomo sapiens]	2e-17	
		Lactococcus lactis					
		cremoris plasmid					
		pNZ4000 insertion					
		sequence IS982					
		putative transposase	ł				
		gene and eps gene	.[(U28739) similar to alcohol		
		cluster	Į.		dehydrogenase/ribitol	ł	
		(epsRXABCDEFGHI	1		dehydrogenase [Caenorhabditis	Ì	
904	U93364	JKL), complete cds	0.007	2731377	elegans]	1e-31	
l		Rattus norvegicus	i				
006	A 50000 co	homer-1c mRNA.	j		i	ŀ	
905	AF093268	complete cds	0.006	<none></none>	<none></none>	<none></none>	
		Mus musculus	1				
906	AF100694	Pontin52 mRNA,	0.004			ĺ	
~~	AF 100094	complete cds	0.006	<none></none>	<none></none>	<none></none>	
ł		Sambucus nigra	ŀ				
		hevein-like protein					
907	AF074386	mRNA, complete cds	0.006	-NONE			
-+	. 2 5/4300	madia, complete cas	0.006	<none></none>	<none></none>	<none></none>	
1		Arabidopsis thaliana	i i	į	l	Í	
1		cellulose synthase]				
ŀ		catalytic subunit (Ath-	j		·		
		B) mRNA, complete			1	[
908		cds	0.006	<none></none>	NONE	NONTE	
		Arabidopsis thaliana		SHORES		<none></none>	
		mRNA for	1			J	
		neoxanthin cleavage	1	_		ĺ	
909		enzyme	0.006	<none></none>	<none></none>	<none></none>	
					NIONES .	(11)11E2	

	Nearest	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC		Treignbor (Blastia vs. C	Jenomik)	Nearest Neigh	nbor (BlastX, vs. Non-Redundant P	roteins)		
D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
<u> </u>								
1	ł	Arabidopsis thaliana	J					
1	I	celiulose synthase	Ì	İ		1		
1	1	catalytic subunit (Ath-	-					
1		B) mRNA, complete		1				
910	AF027174	cds	0.006	<none></none>	<none></none>	<none></none>		
		Rattus norvegicus						
911	A E00224	homer-1c mRNA,				1		
911	AF093268	complete cds	0.006	<none></none>	<none></none>	<none></none>		
		Rattus norvegicus						
912	A EDOSOCI	homer-1c mRNA.			1			
912	AF093268	complete cds	0.006	· <none></none>	<none></none>	<none></none>		
1		Brassica rapa mRNA	٠.					
913	AB012106	for SRK45, complete	0.006	NO.				
1,13	AD012100	cus	0.006	<none></none>	<none></none>	<none></none>		
		Helianthus tuberosus						
1	i	lectin I mRNA.				1		
914	AF064029	complete cds	0.006	<none></none>	100.00			
	14 004025	Mus musculus	0.000	KNONES	<none></none>	<none></none>		
		Pontin52 mRNA.				i i		
915	AF100694	complete cds	0.006	<none></none>	<none></none>	<none></none>		
				4.01.62	(AF063866) ORF MSV064	KNONES		
		Rattus norvegicus	i		hypothetical protein			
		homer-ic mRNA,	l		[Melanoplus sanguinipes	i I		
916	AF093268	complete cds	0.006	4 0 49856	lentomopoxvirus?	9.6		
					(Z82070) predicted using	7.0		
	i	ļ	į		Genefinder; similar to Lectin C-			
	ł	İ	[type domain short and long			
					forms (2 domains); cDNA EST			
			1		EMBL:C10633 comes from this			
			I		gene; cDNA EST			
		Mus musculus			EMBL.C12424 comes from this			
A. 7		Pontin52 mRNA,			gene; cDNA EST yk191e7.3			
917	AF100694	complete cds	0.006	3880536	comes from this	7.9		
1	i.	Sambuana ni	1					
- 1		Sambucus nigra	- 1		(Z81552) F56G4.1			
ł		ribosome inactivating	1		[Cuenorhabditis elegans]			
918		PNA complete add	0.006		>gi 3878615 gnl PID e1348240	_ [
	W 017033	nRNA, complete cds	0.006	3877761	(Z83118) F56G4.1	7.5		
	1,	1.sapiens PTPL1	- 1	}	CATIFOCIALE PRECIMENT	j		
	1	nRNA for protein	1		CATHEPSIN E PRECURSOR	j		
919		vrosine phosphatase	0.006		precursor - rabbit >gi 402729	I		
		resine phosphatase	0.000	1168791	(LOS418) procathepsin E	7.4		

Single Park	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (Blast X vs. Non-Redundant Proteins)			
SEQ		(Diast. + VS. C.	,chomix)	Treatest (vergnoof (Blasic VS. 1908-Redundant Proteins)			
ID ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION		
	1	 	1	L ACCESSION	DESCRIPTION	P VALUE	
	 -				KINASE, BETA	 	
1	l		İ		DIACYLGLYCEROL	1	
1	į.		1		KINASE)	ĺ	
İ	i				>gi 477059 pir A47744	1	
1	ſ	Sambucus nigra	ļ		diacylglycerol kinase (EC		
1		hevein-like protein			2.7.1.107) beta - rat 90kDa-		
920	AF074386	mRNA, complete cds	0.006	1346371	diacylglycerol kinase [Rattus		
	1 4 4 7 1 1 5 5	mad vivi complete cus	0.000	1340371	diacyigiveeror kinase (Rattus	5.5	
1		Lycopersicon					
i	1	esculentum class II			l .	ŀ	
		small heat shock		,			
		protein Le-HSP17.6			(D88588) lipoprotein		
921	U72396	mRNA, complete cds	0.006	2196567	[Escherichia coli]	4.3	
ł	_	Sambucus nigra]	
		hevein-like protein			(Z83259) AmphiBrf38	1 1	
922	AF074387	mRNA, complete cds	0.006	2113798	[Branchiostoma floridae]	4.3	
		Brassica rapa mRNA					
022	ABOLOLOG	for SRK45, complete			(U58282) Bowel [Drosophila		
923	AB012106	cds	0.006	1388166	melanogaster]	4.3	
		Sambucus nigra	1				
		hevein-like protein	ŀ		HYPOTHETICAL 20.1 KD		
924	AF074386	mRNA, complete cds	0.006	2496785	PROTEIN Y4YS	4.2	
					A-AGGLUTININ		
					ATTACHMENT SUBUNIT		
		1	j		PRECURSOR	!	
		Sambucus nigra	ł		>gi 101170 pir A41258 a-		
		ribosome inactivating	1		agglutinin core protein AGA1 -		
		protein precursor	İ		yeast (Saccharomyces		
925	AF012899	mRNA, complete cds	0.006	416592 -	cerevisiae)	2 .7 ·	
.			. 1		A-AGGLUTININ		
j	· 1		ł		ATTACHMENT SUBUNIT	j	
			j		PRECURSOR		
į	1	Halianak			>gi 101170 pir A41258 a-	- 1	
ł		Helianthus tuberosus	į		agglutinin core protein AGA1 -	- !	
926		lectin 1 mRNA,	0.004	414500	veast (Saccharomyces		
/20		Complete cds Arabidopsis thaliana	0.006	416592	cerevisiae) (U41263) The 3' UTR of this	2.5	
		mRNA for	1		gene overlaps the 3' UTR of	1	
1		neoxanthin cleavage	1		T19D12.6(confirmed by EST	ł	
927		enzvme	0.006	3258584	hits) [Caenorhabditis elegans]	2.0	
		Human Down	- 5.555	2230304	mas (Caciomabulus elegans)	2.0	
ł		Syndrome region of	į				
j		chromosome 21.	-			l	
1	Í	genomic sequence.	1		(AF067150) beta-hydroxyacyl-	į	
928		lone A12H1-1A6.	0.006	3850997	ACP dehydratase precursor	1.9	

	Negrest	Neighbor (BlastN vs. C	Panhank)	None No	H. O. W. N. D.	
SEQ		Total Plastit VS. C	i i i i i i i i i i i i i i i i i i i	ivearest iveig	hbor (BlastX vs. Non-Redundant l	roteins)
ID ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-		A 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				
		Arabidopsis thaliana cellulose synthase catalytic subunit (Ath-			·	
1175	AF027173	A) mRNA, complete cds	2e-04	<none></none>	<none></none>	<none></none>
1176	Y09232	H.sapiens fertilin alpha pseudogene	2e-04	<none></none>	<none></none>	<none></none>
1177	AJ005813	Arabidopsis thaliana mRNA for neoxanthin cleavage enzyme	2e-04			
1	73003813	Mus musculus	26-04	<none></none>	<none></none>	<none></none>
1178	AF100694	Pontin52 mRNA,	2e-04	<none></none>	<none></none>	<none></none>
1179	AF072847	Homo sapiens putative swelling- activated chloride channel (CLNS1A) gene, intron 6	2e-04	<none></none>	<none></none>	<none></none>
1180		Sambucus nigra ribosome inactivating protein precursor mRNA, complete cds	2e-04	<none></none>	<none></none>	<none></none>
1181		Sambucus nigra ribosome inactivating protein precursor mRNA, complete cds	2e-04	<none></none>	NONE	NOVE
		Arabidopsis thaliana cellulose synthase catalytic subunit (Ath-A) mRNA, complete	20-04	NOINE	<none> (U50199) coded for by C. elegans cDNA yk89e9.5; coded for by C. elegans cDNA cm7g5; coded for by C. elegans cDNA cm14b9; coded for by C. elegans cDNA yk52g5.5; coded for by C. elegans cDNA yk76e5.5; coded for by C.</none>	<none></none>
1182		ds	2e-04	1213557	elegans cDNA yk131f11.5; c	8.4

	Neares	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC			T T	ivearest iver	enbor (BlastX Vs. Non-Redundant Pi	roteins)		
Ð	- 1	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
<u> </u>					EPITHELIAL DISCOIDIN	1		
- [1	1	1		DOMAIN RECEPTOR I	 		
1	I	İ	1		PRECURSOR (TYROSINE-	ľ		
- [1	I	1	1	PROTEIN KINASE CAK)	l		
1	1	Lycopersicon	i	1	(CELL ADHESION KINASE)	i .		
1	ł	esculentum cytosolic	1	} ·	(TYROSINE KINASE DDR)	1		
į.	1	class II small heat	ĺ	i	(DISCOIDIN RECEPTOR	j 1		
1	1	shock protein HCT2	I	ł	TYROSINE KINASE) (TRK E)			
1183	AF090115	(HSP17.4) mRNA,		1	(PROTEIN-TYROSINE			
1103	WL030112	complete cds	2e-04	729008	KINASE RTK 6) sapiens]	8.3		
	ł	1		Í	HYPOTHETICAL 138.1 KD			
1	l	Sambucus nigra			PROTEIN IN MOLR-BGLX			
1	ł	1	1]	INTERGENIC REGION			
1		ribosome inactivating			>gi 1788436 (AE000300)			
1184	AF012899	protein precursor mRNA, complete cds	2 24		putative regulator (Escherichia	i		
1.04	A1 012099	mikina, complete cas	2e-04	2507582	coli]	7.8		
1	İ	Ì			collagen alpha 1(1X) chain -			
1	l	Sambucus nigra			mouse musculus]	l i		
		hevein-like protein			>gi 744962 prf 2015346A	ł		
1185	AF074386	mRNA, complete cds	2e-04	1005500	collagen:SUBUNIT=alpha1:ISO	I		
		mat v.i. complete cus	26.04	1085500	TYPE=IX [Mus musculus]	7.8		
		Arabidopsis thaliana			1	i		
i l	j	cellulose synthase			1	j		
1		catalytic subunit (Ath-			1	i		
		A) mRNA, complete			(V13042) GTN D-1	1		
1186		cds	2e-04	2623967	(Y13942) GTN Reductase			
				2023907	[Agrobacterium radiobacter]	7.4		
1			i		GLYCOSYLATION END	į		
1			I		PRODUCT-SPECIFIC	ı		
j	Ī	1	j		RECEPTOR PRECURSOR	ì		
	Į	1	j		(RECEPTOR FOR			
	l	· 1			ADVANCED	1		
' I	• 1		· 1		GLYCOSYLATION END			
. 1	1		1.		PRODUCTS) products receptor	1		
ı	-	Arabidopsis thaliana	1		precursor - bovine >gi 163651	- 1		
- 1	ļr.	nRNA for	- 1		(M91212) receptor for advanced	1		
]		neoxanthin cleavage	1		glycosylation end products [Bos	1		
1187	AJ005813 e	enzyme	2e-04	2497316	taurus]	5.3		
- 1	i.							
1		Arabidopsis thaliana	1			j		
]		ellulose synthase	1	•		1		
ľ		atalytic subunit (Ath-	1					
1188		B) mRNA, complete	1					
	AF04/1/4 C	ds	2e-04	1001710	(D64004) hypothetical protein	3.5		

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		July 100 (Diabate 15)		- rearest regin	Doi (Diase 2 73: 110)1-120dundani P	oteins)	
ID	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Arabidopsis thaliana			(U41263) The 3' UTR of this		
		mRNA for			gene overlaps the 3' UTR of		
1		neoxanthin cleavage			T19D12.6(confirmed by EST	1	
1189	AJ005813	enzyme	2e-04	3258584	hits) [Caenorhabditis elegans]	2.1	
		Arabidopsis thaliana			İ		
1		cellulose synthase				;	
1 1		catalytic subunit (Ath-		,		1	
1190	AF027173	A) mRNA, complete	2 04	077/770	(AF038623) contains similarity		
1190	AF02/1/3	cds	2e-04	2736338	to RNA recognition motifs	0.89	
		Lycopersicon					
		esculentum class II		٠,		· .	
		small heat shock				ļ	
1		protein Le-HSP17.6			(D88588) lipoprotein		
1191	U72396	mRNA. complete cds	2e-04	2196567	(Escherichia coli)	0.69	
1	0,2370	micryA. complete cos	20-04	2190307	[[ESCHETICHIA CON]	0.09	
		Lycopersicon					
1 1		esculentum cytosolic]	
1 1		class II small heat				1	
		shock protein HCT2					
1 1		(HSP17.4) mRNA.	-		(AJ006096) F-spondin		
1192	AF090115	complete cds	2e-04	33 19874	[Branchiostoma floridae]	5e-04	
		Chlamydomonas	1		1		
		reinhardtii dynein					
		heavy chain alpha	į		(Z81077) predicted using		
	1.000.0	(ODA11) gene, exons			Genefinder: Similarity to Yeast		
1193	L26049	2-15, and partial cds.	2e-04	3876775	protein 8248 (TR:G587531)	2e-09	
1 1		Mus musculus Pontin52 mRNA.	ŀ				
1194	AF100694		1e-04	ANONES	NONE	ANONIT:	
1194	AT 100094	complete cds	16-04	<none></none>	<none></none>	<none></none>	
1		Helianthus tuberosus	ſ				
[lectin 1 mRNA.	· [·		
1195	1	complete cds	1c-04	<none></none>	<none></none>	<none></none>	
		Tomprete eas	-15.54	WI OTTES	3000	~!!O!!!	
1	j	Homo sapiens]				
		retinaldehyde-binding]				
		protein (CRALBP)	1			.	
1196	L34219	gene, complete cds.	1c-04	<none></none>	<none></none>	<none></none>	
		Rhesus monkey					
1197	X51890	interleukin-3 gene	1e-04	<none></none>	<none></none>	<none></none>	

	Nearest I	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ						T	
D D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Plasmodium				 	
		falciparum				· 	
		chromosome 2.		· ·	}		
		section 58 of 73 of					
		the complete		i –	}]	
1198	AE001421	sequence	le-04	<none></none>	<none></none>	<none></none>	
1 1		Lycopersicon			ļ		
		esculentum cytosolic					
[class II small heat				1	
		shock protein HCT2				1	
1100	AFOOOLIE	(HSP17.4) mRNA.	1- 04	, NONE.	Nove		
1199	AF090115	complete cds	1e-04	<none></none>	<none></none>	<none></none>	
1 1		Arabidopsis thaliana	• .				
		cellulose synthase					
		catalytic subunit (Ath-	-			1	
		B) mRNA, complete			(Y15086) HepC protein		
1200	AF027174	cds	1e-04	2576287	[Cylindrotheca fusiformis]	4.7	
		Arabidopsis thaliana			[-7,1		
		mRNA for					
		neoxanthin cleavage			(AB016623) RWC-3 [Oryza	Ĭ,	
1201	AJ005813	enzvme	1e-04	3395673	satival	0.14	
		Homo sapiens					
		BRCA1-associated				1 1	
		RING domain protein					
		(BARDI) gene,			i .		
1202		exons 2 and 3	9e-05	<none></none>	<none></none>	<none></none>	
		Arabidopsis thaliana				0.0	
1	1	mRNA for					
		neoxanthin cleavage					
1203		enzyme	9e-05	<none></none>	<none></none>	<none></none>	
		Brassica rapa mRNA	1			[i	
1		for SRK45, complete	· <u> </u>	•	ľ		
1204		cds	9e-05	<none></none>	<none></none>	<none></none>	
1		Xenopus laevis	i		l	 	
- 1		mitotic	1	•]		
1205		phosphoprotein 44	0.05	NONE			
1205	U95098	mRNA, partial eds	9e-05	<none></none>	<none> HYPOTHETICAL</none>	<none></none>	
- 1]		1		LIPOPROTEIN MG348		
	l				PRECURSOR		
	[,	Laurania higglas	1		PRECURSOR >gi 1361668 pir E64238		
- 1		Laccaria bicolor	ı				
		glyoxal malate			hypothetical protein MG348 -		
1206		synthase protein	02.05	1251552	Mycoplasma genitalium (SGC3)		
1200	AF034099	mRNA, complete cds	9e-05	1351553	>gi 3844931	8.8	

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	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Human DNA for					
		alpha-platelet-derived			(AF056494) NADH		
		growth factor			dehydrogenase subunit 5		
1207	D50006	receptor, exon 6-10	9e-05	3063639	[Panorpa japonica] INHIBIN ALPHA CHAIN	5.1	
		Human Down					
		Syndrome region of	I	•	PRECURSOR bovine		
		chromosome 21.		10.4077	>gi 163195 (M13273) inhibin A	20	
1208	U50423	clone A41B8-1B7.	9e-05	124273	subunit [Bos taurus] (X72850) 2.4-	3.0	
		Arabidopsis thaliana			dihydroxybenzoate		
		mRNA for			monooxygenase [Sphingomonas	1	
		neoxanthin cleavage	0-05	4007780		2.3	
1209	AJ005813	enzyme Homo sapiens clone	9e-05	4007782	sp.]	2.3	
		fragment			1	•	
}		UWGC:gap3 from			1		
		7q31.3, complete				1	
		sequence [Homo			(U60315) MC132L [Molluscum		
1210	AC005276	sapiens]	9e-05	1492075	contagiosum virus subtype []	1.0	
1210	AC003270	Mus musculus	- 20 03	1472073	Contagios on the sacty por ty	1.0	
		Pontin52 mRNA.			(AB007884) KIAA0424 [Homo		
1211	AF100694	complete cds	9e-05	2887423	sapiens]	2e-10	
	12 1000)				(U83115) non-lens beta gamma-		
		C.fuscus gamma-M2-			crystallin like protein [Homo		
1212	X77772	1 crystallin mRNA.	9e-05	2072425	sapiens]	7e-25	
		Brassica rapa mRNA					
		for SRK45, complete	1				
1213	AB012106	cds	8e-05	<none></none>	<none></none>	<none></none>	
		Apis mellitera		•			
		ligustica complete					
		mitochondrial					
1214	L06178	genome	8e-05	<none></none>	<none></none>	<none></none>	
		Brassica rapa mRNA				, , , , , ,	
		for SRK45, complete			11017	AVONT-	
1215	AB012106	cds	8e-05	<none></none>	<none></none>	<none></none>	
		C 1					
		Sambucus nigra	ŀ				
		ribosome inactivating					
اء.ي ا	1176504	protein precursor	8e-05	<none></none>	<none></none>	<none></none>	
1216	U76524	mRNA, complete cds Apis mellifera	96-03	SINUNES	CINONE	VIIOI17	
		Apis mentiera ligustica complete					
ł		mitochondrial	- 1				
1217		mitocnondriai	8e-05	<none></none>	<none></none>	<none></none>	
121/	L06178	Brassica rapa mRNA	00-05	VI OILL			
1		for SRK45, complete	1				
1218		cds	8e-05	<none></none>	<none></none>	<none></none>	
1-10	YP017100	- us	00-05	2,101127	1 3.101122		

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEO							
ID ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Mus musculus		7.002.00.01		1	
		Pontin52 mRNA.				 	
1219	AF100694	complete cds	8e-05	<none></none>	<none></none>	<none></none>	
F	111 10007	Brassica rapa mRNA	-33 32			VIII OILE	
		for SRK45, complete					
1220	AB012106	cds	8e-05	<none></none>	<none></none>	<none></none>	
					WNT-11 PROTEIN		
		Brassica rapa mRNA	İ		PRECURSOR (XWNT-11)	ĺ	
		for SRK45, complete			clawed frog >gi 439108	ł	
1221	AB012106	cds	8e-05	1722841	(L23542) maternal protein	9.9	
		Arabidopsis thaliana					
		cellulose synthase					
		catalytic subunit (Ath-					
		B) mRNA, complete	1		(U35637) nebulin (Homo		
1222	AF027174	cds	8e-05	1205991	sapiens]	9.6	
			ı		·		
		Homo sapiens serine					
		protease-like protease					
		Sequence 2 from			(AF055354) respiratory burst		
1223		patent US 5736377	8e-05	3242783	oxidase protein B	8.6	
		Rattus norvegicus	·				
		mRNA for PAG608		22.12.2	(AE000616) alpha-ketoglutarate		
1224	Y13148	gene	8e-05	2314243	permease (kgtP) RAS GTPASE-ACTIVATING-	8.1	
i I					LIKE PROTEIN IQGAPI		
					(P195) (KIAA0051)		
1 1	.				>gi 627594 pir A54854 Ras		
					GTPase activating-related		
li	İ	Arabidopsis thaliana	1		protein - human sapiens]		
		mRNA for	ĺ		>gi 536844 (L33075) ras		
i	ľ	neoxanthin cleavage	J		GTPase-activating-like protein		
1225		enzyme	8e-05	1170586	[Homo sapiens]	7.8	
					[
	İ		1	•	NADH-UBIQUINONE		
l		1	1		OXIDOREDUCTASE CHAIN		
	ļ		j		4 >gi 1085185 pir S52968		
1	1	Arabidopsis thaliana	1		NADH dehydrogenase chain 4 -		
	ļ	cellulose synthase	1		honeybee mitochondrion		
	į,	catalytic subunit (Ath-	1		(SGC4) >gi 552446 (L06178)		
I	ļ.	A) mRNA, complete	i	•	NADH dehydrogenase subunit 4	j	
1226	1	cds	8 c -05	464239	[Apis mellifera ligustica]	3.5	
		Mus musculus					
		Pontin52 mRNA.	Ì				
1227	AF100694	complete cds	8e-05	544353	F-SPONDIN PRECURSOR	3.5	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Sambucus nigra					
j		ribosome inactivating				1	
1 1		protein precursor			apolipoprotein B-100 - chicken]	
1228	U76524	mRNA, complete cds	8e-05	483243	(fragment)	3.4	
		Rattus norvegicus					
		homer-1c mRNA.			proline-rich protein - mouse		
1229	AF093268	complete cds	8e-05	91207	(fragment) musculus]	2.2	
		Arabidopsis thaliana					
		cellulose synthase					
1 1		catalytic subunit (Ath-					
1 1		A) mRNA, complete		• •	ZONADHESIN PRECURSOR		
1230	AF027173	cds	8e-05 [.]	2499181	>gi 1066466	2.2	
1230	AF02/1/3	cus	80-05	2499101	Zgi 1000400	2.2	
		Arabidopsis thaliana					
ļ		cellulose synthase				[
		catalytic subunit (Ath-					
		A) mRNA, complete			ZONADHESIN PRECURSOR		
1231	AF027173	cds	8e-05	2499181	>gi 1066466	1.9	
		Brassica rapa mRNA					
		for SRK45, complete			(AF027972) flagelliform silk	1	
1232		cds	8e-05	2833647	protein [Nephila clavipes]	1.6	
		Rattus norvegicus					
		homer-1c mRNA,	[(Z49821) MYO2	0.00	
1233	AF093268	complete cds	8e-05	1163063	[Saccharomyces cerevisiae]	0.90	
		A bidansis shaliana					
		Arabidopsis thaliana					
!		cellulose synthase catalytic subunit (Ath-					
		B) mRNA, complete					
1234	AF027174	cds	8e-05	1653488	(D90914) hypothetical protein	0.30	
1234	A102/1/4	Chicken nonmuscle	86-03	1023400	1030314) hypothetical protein	5.50	
		myosin heavy chain	ŀ	•			
		(MHC) gene,]	•	1		
1235	M26510	complete cds.	8e-05	112159	plectin - rat	0.003	
1223		Human chromatin					
		structural protein			(AF003384) weak similarity to		
1236		homolog	8e-05	2088823	the peptidase family A2	1e-13	
		Mus musculus					
		Pontin52 mRNA.			(U02289) GTPase-activating		
1237		complete cds	8e-05	437181	protein [Caenorhabditis elegans]	2e-17	
		Mus musculus			HYPOTHETICAL 80.8 KD		
	1	Pontin52 mRNA.	i		PROTEIN ZC21.4 IN		
1238	AF100694	complete cds	8e-05	465983	CHROMOSOME III	8e-27	

	Nearest N	leighbor (BlastN vs. G	enbank)	Nearest Neight	oor (BlastX vs. Non-Redundant Pr	oteins)
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
1239	AF090115	Lycopersicon esculentum cytosolic class II small heat shock protein HCT2 (HSP17.4) mRNA, complete cds	7e-05	<none></none>	. <none></none>	<none></none>
		Rattus norvegicus NF- KB gene, promotor			(AL031633) predicted using Genefinder; cDNA EST yk304f12.5 comes from this gene [Caenorhabditis elegans]	9.3
1240	U83656 AF074387	Sambucus nigra hevein-like protein mRNA, complete cds	7e-05 7e-05	3880858	(AL022600) hypothetical	9.2
1242	X89398	H.sapiens ung gene for uracil DNA- glycosvlase	7e-05	549700	HYPOTHETICAL 23.7 KD PROTEIN IN MDH1-VMA5 INTERGENIC REGION >gi 539182 pir S37908 hypothetical protein YKL083w - yeast (Saccharomyces cerevisiae) >gi 486120 (Z28082) ORF YKL083w	1.8
1243	M83753	Bovine follicle stimulating hormone- beta subunit gene. complete cds.	7e-05	2398621	(AJ000342) DMBT1 protein, 5.8 kb transcript [Homo sapiens]	1.8
1244	M80829	Rat troponin T cardiac isoform gene, complete cds	5e-05	854065	(X83413) U88 [Human herpesvirus 6]	2e-08
1245	AF074387	Sambucus nigra hevein-like protein mRNA, complete cds	4e-05	120240	FLAGELLIN B2 PRECURSOR Methanococcus voltae >gi 150063 (M72148) flagellin	5.2
1246	AF012899	Sambucus nigra ribosome inactivating protein precursor mRNA, complete cds	3e-05	<none></none>	<none></none>	<none></none>
1247		Sambucus nigra ribosome inactivating protein precursor mRNA, complete eds	3e-05	<none></none>	<none></none>	<none></none>
1248	AF074386	Sambucus nigra hevein-like protein mRNA, complete cds	3e-05	<none></none>	<none></none>	<none></none>

PCT/US00/18374

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	Nearest I	verginoi (Brasil Vs. Ci	citoanicy	Treatest Freight	1	l l	
SEQ		DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID	ACCESSION		FVALUE	ACCESSION) Description	T VALUE	
		Rattus norvegicus					
1010	A F003340	homer-1c mRNA.	3e-05	<none></none>	<none></none>	<none></none>	
1249		complete cds Brassica rapa mRNA	30-03	ZITOTIES_	1.0	4.0.,2	
		for SRK45, complete			(AF039716) Similar to protein		
1250	AB012106	cds	3e-05	2773226	kinase (Caenorhabditis elegans)	6.7	
1.230	7.2012100	Mus musculus					
		Pontin52 mRNA,			(U93568) putative p150 [Homo		
1251	AF100694	complete cds	3e-05	2072961	sapiens]	5.6	
					EXOGLUCANASE II	[
					PRECURSOR cellulose 1.4-beta cellobiosidase (EC 3.2.1.91) II	1	
				**	precursor - fungus (Trichoderma	·	
		Lycopersicon			reesei) 1,4-beta-cellobiosidase		
		esculentum class II small heat shock	"		(EC 3.2.1.91) II - fungus		
		•			cellobiohydrolase II		
1252	1172206	protein Le-HSP17.6 mRNA, complete cds	3e-05	121855	[Trichoderma reesei]	4.6	
1252	<u>U72396</u>	mkna, complete cus	36-03	121033	(Trichoderma (eess)		
1	:	Sambucus nigra			(AL021572) similar to CTP		
		ribosome inactivating			SYNTHASE (EC 6.3.4.2) (UTP-	4	
		protein precursor			-AMMONIA LIGASE) (CTP		
1253	U76524	mRNA, complete cds	3e-05	3880516	SYNTHETASE)	3.3	
		Mouse brain-1 POU-					
		domain protein,			(U66102) intimin [Escherichia	3.0	
1254	M88299	complete cds.	3e-05	1947048	coli] CELL-CYCLE NUCLEAR	3.0	
					AUTOANTIGEN SG2NA		
		Vanana la suis			(S/G2 NUCLEAR ANTIGEN)		
		Xenopus laevis			>gi 1082650 pir JC2522 nuclear		
}	4	phosphoprotein 44			autoantigen - human >gi 305095		
1255	U95098	mRNA, partial cds	3e-05	3122872	(U17989) GS2NA	2.8	
1223	073070	matter, partial cas			CYTOCHROME C OXIDASE		
}					POLYPEPTIDE I chain I -] .	
		Sambucus nigra			Thermus aquaticus >gi 155083		
]]		ribosome inactivating			(M84341) cytochrome c oxidase]	
		protein precursor			subunits precursor [Thermus		
1256	U76524	mRNA, complete cds	3e-05	1352145	thermophilus]	2.6	
					SEGMENTATION POLARITY		
		Lycopersicon			PROTEIN ENGRAILED >gi]2076747 (U-2-129)		
		esculentum class II			engrailed (Anopheles gambiae)		
}		small heat shock			>gi 2148918 (U42214)		
ا ا	117000	protein Le-HSP17.6	30.05	2811015	engrailed (Anopheles gumbiae)	2.0	
1257	U72396	mRNA, complete cds	3e-05	2011013	Tensianen (Amobiletes Samoine)		

	Nanzect N	Neighbor (BlastN vs. Ge	nhank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	[4earest t	tergribor (Blastiv vs. O	nounk,	1.02.000 1.0101			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Arabidopsis thaliana	·				
		cellulose synthase	Ī		•		
		catalytic subunit (Ath-			(U62325) FE65-like protein		
		B) mRNA, complete	2.06	1657 7 52	[Homo sapiens]	1.7	
1258	AF027174	cds Mus musculus	3e-05	103/732	[Homo sapiens]	***	
		Pontin52 mRNA.			(U93568) putative p150 [Homo		
1259	AF100694	complete cds	3e-05	2072961	sapiens	1.5	
1209	AF100094	Complete cus	- 30 03	30.2701	CYTOCHROME C OXIDASE		
		ļ			POLYPEPTIDE I chain I -	i .	
		i			Thermus aquaticus >gi 155083		
		Sambucus nigra lectin			(M84341) cytochrome c oxidase		
		precursor mRNA.			subunits precursor [Thermus		
1260	U76523	complete cds	3e-05	1352145	thermophilus]	1.1	
		H.sapiens regulatory			Sxr (Bkm-homolog) sex-		
		region of HOXA7			determining region protein -		
1261	X91890	gene	3e-05	111013	mouse	1.0	
					(D84239) IgG Fc binding		
	* 24224	Homo sapiens metase	3e-05	1944352	protein [Homo sapiens]	0.99	
1262	L36936	gene, partial cds	36.03	1944332	protetti [t tomo sapiens]	3,77	
1	ĺ		·		SMP2 PROTEIN		
					>gi 320853 pir S30911 SMP2		
		Brassica rapa mRNA			protein - yeast (Saccharomyces		
		for SLG45, complete			cerevisiae) gene		
1263	AB012105	cds	3e-05	- 417782	[Saccharomyces cerevisiae]	0.89	
						ļ	
		Sambucus nigra			INTEGRIN ALPHA CHAIN-		
		ribosome inactivating		•	LIKE PROTEIN alpha Intip		
,,,,		protein precursor	3e-05	1708501	[Candida albicans]	0.39	
1264	U76524	mRNA, complete cds	36.03	1700301	(Canada dioicans)		
		Lycopersicon		•			
		esculentum cytosolic					
		class II small heat					
		shock protein HCT2					
		(HSP17.4) mRNA.			cis-Golgi matrix protein GM130		
1265	AF090115	complete cds	3e-05	1587031	[Rattus norvegicus]	0.20	
		Human DNA					
		sequence from					
		cosmid U65A4,			1		
		between markers			(U93569) putative p150 [Homo		
1266	Z81014	DXS366 and DXS87 on chromosome X *	3e-05	2072964	supiens)	0.049	
1.200	Z01U14	Iou chiomosome A	30.03	-0204	1		

	Nearest	Neighbor (BlastN vs. C	ienbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ				- Total est inergi	Tool (Blasce 13: 110) Redundant F	Totellis)	
ID	ACCESSION	DESCRIPTION	DVALUE	A CORDONAL	DESCRIPTION.		
	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
	ļ	 			glycosylated and mynstilated		
1	Ì	ĺ			smaller surface antigen -		
	1		l		Plasmodium falciparum		
1	f	**		İ	>gi 836640 (X76298)	1	
		H.sapiens telomeric	ŀ		glycosylated and myristilated	İ	
1		DNA sequence, clone			smaller surface antigen gallus]	1	
		7PTEL001, read			>gi 1092178 prf 2023165B	i	
1267	Z96668	7PTELOO001.seq	3e-05	542429	surface antigen	0.029	
					(Z70310) predicted using		
		1			Genefinder; Similarity to Mouse	J	
i					ankyrin (PIR Acc. No. S37771);		
	ł			· ·	cDNA EST EMBL:T01923	ĺ	
ĺ					comes from this gene; cDNA		
1	ł				EST EMBL:D32335 comes		
					from this gene; cDNA EST	[]	
					EMBL:D32723 comes from this		
			İ		gene; cDNA ES Genefinder:		
				•	15		
					Similarity to Mouse ankyrin		
					(PIR Acc. No. \$37771); cDNA		
			1		EST EMBL:T01923 comes		
			j	•	from this gene; cDNA EST		
		Brassica rapa mRNA			EMBL:D32335 comes from this		
		for SLG45, complete	ľ		gene; cDNA EST		
1268	AB012105	cds	3e-05	2070121	EMBL:D32723 comes from this		
1200	AD012103	cus	3e-03	3879121	gene; cDNA ES	2e-13	
		Sambucus nigra			ZYXIN (ZYXIN 2) sapiens]	1	
ı		hevein-like protein	1		>gi 1545954 gnl PID e223417		
1269	AF074385	mRNA, complete cds	3e-05	2497677	(X95735) zyxin	2e-23	
				2-77077	(A22733) 2VAIII	26-23	
	ļ	Arabidopsis thaliana	ł				
		cellulose synthase	1			ł	
j		catalytic subunit (Ath-	·	•			
İ		B) mRNA, complete	1				
1270		cds	le-05	<none></none>	<none></none>	<none></none>	
I					PITUITARY HOMEOBOX 3		
	ŀ		1		(HOMEOBOX PROTEIN	i	
1		Canine mRNA for	l		PITX3) >gi 2645427		
		signal recognition			(AF005772) homeobox protein		
1271		particle 54k protein	I e ∙05	3122612	Pitx3 (Mus musculus)	4.4	
- 1		Brassica rapa mRNA	7		(D90905) DNA mismatch repair		
[for SLG45, complete			protein MutL [Synechocystis	ļ	
1272	AB012105	eds ·	1e-05	1652458	sp.]	0.62	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
1273	U57843	Human phosphatidylinositol 3-kinase delta catalytic subunit mRNA, complete cds	le-05	475909	(X67098) ORFIA [Homo sapiens]	0.22	
1274	Z 96569	H. sapiens telomeric DNA sequence, clone 2QTEL054, read 2QTEL00054.seq	le-05	2137043	unknown protein - rabbit (fragment) cuniculus]	0.005	
		Methanobacterium thermoautotrophicum from bases 172512 to 182957 (section 16 of 148) of the complete			kinensin-like protein KIF4 (SW:P33174); cDNA EST EMBL:D27320 comes from this gene; cDNA EST EMBL:D27322 comes from this gene; cDNA EST EMBL:D27321 comes from this gene; cDNA EST EMBL:D35764 comes Mouse kinensin-like protein KIF4 (SW:P33174); cDNA EST EMBL:D27320 comes from this gene; cDNA EST EMBL:D27322 comes from this gene; cDNA EST EMBL:D27322 comes from this gene; cDNA EST EMBL:D27321 comes from this gene; cDNA EST		
1275	AE000810	genome	1e-05	3877579	EMBL:D35764 comes	6e-27	
1276	AB012113	Homo sapiens gene for CC chemokine PARC precursor, complete cds	9e-06	<none></none>	<none></none>	<none></none>	
1210		Homo sapiens Xp22- 154-155 BAC GSHB- 52411 (Genome Systems Human BAC Library), complete sequence (Homo	70-00	310112			
1277		sapiens]	9e-06	<none></none>	<none></none>	<none></none>	
1278	D86245	Human MHC (HLA) DRB intron 1 DNA, partial sequence	9e-06	1051253	(U37531) mucin apoprotein [Mus musculus]	1.3	
1279	D79998	Human mRNA for KIAA0176 gene. partial cds	9e-06	2833253	HYPOTHETICAL PROTEIN KIAA0176 sapiens]	4e- 0 6	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
					(269655) Similarity to Yeast		
					uridine kinase		
					(SW:URK1_YEAST); cDNA		
ļ					EST EMBL:Z14695 comes		
				ı	from this gene; cDNA EST		
		Toxoplasma gondii			CEMSE17F comes from this		
		RH uracil			gene; cDNA EST	ł i	
		phosphoribosyl			EMBL:D67355 comes from this		
		transferase gene.			gene; cDNA EST yk209h1.5		
1280	U10246	complete cds.	9e-06	3876090	comes from this ge	7e-33	
i i					(Z69635) Similarity to Yeast		
I I		• •		• •	uridine kinase		
1 1					(SW:URK1_YEAST); cDNA EST EMBL:Z14695 comes	1	
			. ,		from this gene; cDNA EST		
		Tauadaana aandii			CEMSE17F comes from this		
		Toxoplasma gondii RH uracil			gene; cDNA EST		
		phosphoribosyl			EMBL:D67355 comes from this		
		transferase gene,			gene; cDNA EST yk209h1.5		
1281	U10246	complete cds.	9e-06	3876090	comes from this ge	7e-34	
1201	010240	complete cus.	36-00	3670030	comes from this go	70-54	
1 I		Sambucus nigra			·	i	
		ribosome inactivating				1	
		protein precursor					
1282	AF012899	mRNA, complete cds	8e-06	<none></none>	<none></none>	<none></none>	
		Sambucus nigra					
		ribosome inactivating					
		protein precursor					
1283		mRNA, complete cds	8e-06	<none></none>	<none></none>	<none></none>	
	111111111	Human Rh blood					
		group C antigen					
		(RHCE) gene, exon			(U80837) F07E5.6 gene product		
1284	U66340	2. partial cds	8e-06_	17.07 155	[Caenorhabditis elegans]	9.6	
		Sambucus nigra					
		ribosome inactivating			1.0		
		protein precursor					
1285	AE012800		7e-06	<none></none>	<none></none>	<none></none>	
1202	AF012899	mRNA, complete cds Human insulin		(110112)	3.10.132		
	j	receptor (allele 2)	·				
		gene. exons 14, 15.					
1286	M29930	16 and 17.	4e-06	<none></none>	. <none></none>	<none></none>	
		Homo sapiens					
		(subclone 5_d3 from			i		
		P1 H25) DNA					
1287	L42103	sequence.	3e-06	<none></none>	<none></none>	<none></none>	

	Nearest	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ				Total Total				
ΙD	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION			
		Mus musculus	T TIEGE	ACCESSION	DESCRIPTION	P VALU		
		cerberus-like (Cer-I)		 	-			
1288	AF012244	gene, complete cds	3e-06	ANONE:				
1200	711 012244	gene, complete cus	36-00	<none></none>	<none></none>	<none></none>		
		Human DNA						
		sequence from						
		cosmid L96F8.			<i>-</i>	İ		
		Huntington's Disease			1	1		
		Region, chromosome						
1289	Z69366	4p16.3 contains EST.	3e-06	<none></none>	<none></none>	<none></none>		
					- ANGINES	- STYOLY		
1		Human DNA						
		sequence from		•	1			
ı		cosmid L96F8.			1			
		Huntington's Disease						
		Region, chromosome						
1290	Z69366	4p16.3 contains EST.	3e-06	<none></none>	<none></none>	<none></none>		
i		H.sapiens	****					
!		chromosome 3			1			
1291	X85232	sequences	3e-06	<none></none>	· <none></none>	<none></none>		
í		Human platelet						
1202		glycoprotein Illa.						
1292		exons 7, 8 and 9.	3e-06	<none></none>	<none></none>	<none></none>		
- 1		Human HepG2 3'	J					
1293		region cDNA, clone hmd2a0t	3. 00	000504	(U33484) ependymin	l i		
1273		Lagothrix lagotricha	3e-06	998296	[Hemiodus sp.]	5.6		
- 1		interphotoreceptor				1 1		
ı	•	retinoid-binding				i 1		
ı		protein (IRBP) gene.	ı					
ł		intron 1, complete	ı		(U71440) polyprotein [Rice	ì		
1294		sequence	3e-06	1613846	tungro spherical virus]	5.0		
				10150.0	angre spherical virus	3.0		
1	· [:	Lycopersicon		•				
J	ŀ	esculentum cytosolic	·					
j		class II small heat	!					
1		shock protein HCT2	- 1		(U53204) plectin [Homo			
1		HSP17.4) mRNA.	j		sapiens] >gi[1477651 (U63610)			
1295	AF090115 (complete cds	3e-06	1477646	plectin [Homo sapiens]	4.0		
1					reverse transcriptase - fruit fly			
		Iomo sapiens B-ATF			reverse transcriptase	į		
1296	AF016898	ene, complete cds	3e-06	1085177	[Drosophila yakuba]	3.0		
	ŀ	į.	[(Z81522) predicted using			
	١.	, , , , , , , , , , , , , , , , , , , ,	1		Genefinder; similar to RNA	1		
		domo sapiens DNA,	i i		recognition motif. (aka RRM,			
207		rinucleotide repeats	2.01	2004	RBD, or RNP domain)			
297	AB018490 r	egion	3e-06	3876572	[Caenorhabditis elegans]	3.0		

				Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	Nearest N	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		`					
		Arabidopsis thaliana					
		cellulose synthase					
		catalytic subunit (Ath-					
		B) mRNA, complete			(AB020631) KIAA0824 protein	2.7	
1298	AF027174	cds	3e-06	4240137	[Homo sapiens]	2.1	
		Homo sapiens					
		adenosine					
1	-	monophosphate			(D90916) thiol:disulfide		
		deaminase 1			interchange protein DsbD		
		(AMPD1) gene,	2.00	1653775	[Synechocystis sp.]	1.7	
1299	M37929	exons 11-12. Homo sapiens	3e-06	. 1023/13	[Syliection vstis sp.]	• • • • • • • • • • • • • • • • • • • •	
		adenosine			1		
		monophosphate					
İ		deaminase 1			(D90916) thiol:disulfide		
		(AMPDi) gene.			interchange protein DsbD		
1300	M37929	exons 11-12.	3e-06	1653775	[Synechocystis sp.]	1.7	
1300	W137929	Glycine max actin	30 00	1002110	ACTIN-LIKE PROTEIN ARPS		
		(Soy86) gene. partial			Ynl2430p [Saccharomyces		
1301	U60496	cds	3e-06	1730738	cerevisiae	2e-05	
1.301	000470	Yersinia					
,		pseudotuberculosis				İ	
		rplC, rplD, rplW,				1	
1		rplB and rpsS genes	1				
	ļ	for ribosomal proteins			50S RIBOSOMAL PROTEIN	ì	
		L3, L4, L23, L2 and	1		L2 maritima >gi 437926		
1302	X14363	\$19	3e-06_	585879	(Z21677) ribosomal protein L2	2e-12	
		H.sapiens DNA for	[]				
		microsatellite				<none></none>	
1303	Z34969	polymorphism	2e-06	<none></none>	<none></none>	CNOINES	
		H.sapiens BBC1		NOVE	NONES	<none></none>	
1304	X64707	mRNA	1e-06	<none></none>	<none></none>	7110111	
	1	Homo sapiens Xp22-				ļ	
	1	154-155 BAC GSHB-] '		
		524[1 (Genome					
		Systems Human BAC		1	1	Ì	
		Library), complete			1		
1200	4.0005000	sequence [Homo	le-06_	<none></none>	<none></none>	<none></none>	
1305	AC005830	sapiens	10-00	3,101,12			
		Human electron					
		transfer flavoprotein			1		
1		alpha-subunit mRNA.	.]				
1306	J04058	complete cds.	1e-06	<none></none>	<none></none>	<none></none>	
	1 101033	Trombiere res	 	<u> </u>		_	

	Nearest Neighbor (BlustN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
	Necession	J GOOGLE CLOSE		7,0020010.			
		Homo sapiens		-		 	
		fibroblast growth					
		factor receptor gene			1		
		(located in the central					
		MHC) signal peptide		, i	mxcQ gene [Methylobacterium		
1307	L25647	and consecutive exon	le-06	1586734	organophilum]	5.4	
		Human MHC class III			(U20633) NADH		
1308	1 36361		1e-06	1684985	dehydrogenase subunit [Neuwiedia veratrifolia]	1.8	
1308	L26261	HLA-RP1 gene.	12-00	1004763	(Nedwiedia veradifolia)	1.0	
		Mus musculus alpha-		10		[. .	
		actinin-2 associated					
		LIM protein mRNA.			(AF053367) carboxyl terminal		
		alternatively spliced			LIM domain protein [Mus		
1309	AF002283	product, complete cds	le-06	2996196	musculus	4e-17	
		Human haptoglobin					
	•	gene (alpha-2 allele),			·		
		complete cds and haptoglobin-related		•			
		gene, exon 1 and					
1310	M10935	three Alu repeats.	6e-07	<none></none>	<none></none>	<none></none>	
1000		Homo sapiens					
		(subclone l_g6 from					
		BAC H76) DNA			coagulation factor Xa (EC		
1311	AC002251	sequence	4e-07	2144491	3.4.21.6) precursor norvegicus]	4.2	
		C					
! I		Streptomyces chrysomallus					
		actinomycin					
		synthetase II (acmB)	ì		(UI5I81) 4-coumarate-coA	! !	
1312	AF047717	gene, complete cds	4c-07	699196		1e-06	
-					ligase [Mycobacterium leprae] GUANINE NUCLEOTIDE		
			,		DISSOCIATION		
			i		STIMULATOR RALGDS		
					FORM A (RALGEF)		
					>gi 321257 pir S28415 guanine nucleotide dissociation		
		Human Ral guanine		·	stimulator ralGDS - mouse		
		nucleotide dissociation			>gi 193573 (L07924) guanine		
		stimulator mRNA.			nucleotide dissociation		
1313	U14417	partial eds.	4e-07	544402	stimulator [Mus musculus]	8e-08	
1010	014417	H.sapiens flow-sorted		J . / - O Z			
		chromosome 6			1		
		Hindlll fragment.		·			
1314	Z79027	SC6pA20G8	3e-07	<none></none>	<none></none>	<none></none>	

PCT/US00/18374

Homo sap intestinal (MUC2) g	mucin gene. region and	P VALUE	ACCESSION	DESCRIPTION	P VALUE
Homo sap intestinal (MUC2) g promoter 1315 U67167 partial cds	mucin gene. region and	P VALUE	ACCESSION	DESCRIPTION	P VALUE
intestinal (MUC2) g promoter 1315 U67167 partial cds	mucin gene. region and				
(MUC2) g promoter 1315 U67167 partial cds	gene. region and				
promoter 1315 U67167 partial eds	region and		•		
1315 U67167 partial cds	-				1
	:		_		
I Homo can		3e-07	<none></none>	<none></none>	<none></none>
	ert cDNA	3.07	NONE	NONT	<none></none>
1316 AF086256 clone ZD4	ICII	3e-07	<none></none>	<none> (U97003) contains similarity to</none>	<none></none>
1 1				C4-type zinc fingers and a]
Human ch	one HS4.61			ligand-binding domain of	!
1317 U67228 Alu-Ya5 s		3e-07	1938437	nuclear hormone receptors	2.3
1317 007228 Alu-125 s	equence	30-07	1730437	nacion normone receptors	
Human ca	lpain-like	٠.			
protease ((AF047659) No definition line	
	omplete cds	3e-07	2911858	found [Caenorhabditis elegans]	0.39
Homo sap	iens				
SERCA3	gene, exons				
1319 Y15724 1-7 (and j	oined CDS)	le-07	<none></none>	<none></none>	<none></none>
Bean DNA					
, , , , , , , , , , , , , , , , , , , ,	ch cell wall				
1320 X13596 protein GI	RP 1.8	le-07	<none>_</none>	<none></none>	<none></none>
	:				
Homo sap					
	glutathione			*	
peroxidase				(U58751) C07G1.7 gene	
complete				product [Caenorhabditis	
	ne, 3' end.	le-07	1326385	elegans]	8.0
110111111111111111111111111111111111111	, 5 3.10.				
H.sapiens	CpG DNA.	1		· i	
	, forward			extensin-like protein - maize	
1322 Z55905 read cpg7		le-07	1076802	>gi 600118 mays]	0.61
Human m.	RNA of trk				
oncogene				(M74509) [Human endogenous	
gb I96186		Į		retrovirus type C oncovirus	
Sequence		1	000100	sequence.], gene product [Homo	
1323 X03541 patent US	5734039	1e-07	325465	supiens)	3e-04
Canis fam		1		(D10639) sine #========	
linked zing	nnger	10.07	2204.12	(D10628) zinc finger protein [Mus musculus]	7e-08
1324 AF027766 protein Bovine ml	SNA for	le-07	220643	[ivius musculus]	76-08
rabphilin-		İ			
complete		ŀ		(AC004082) rab3 effector-like;	
dbj E0780		1		35% Similarity to AF007836	
cDNA end		ŀ	٠.	(PID:g2317778) [Homo	
1325 D13613 rabphilin-3		le-07	2822161	sapiens]	6e-11

WO 01/02568

	Nearest N	leighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
וס	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Human mRNA for c-			(J04169) gag-one fusion protein		
1326	X57110	cbi proto-oncogene	le-07	323270	[Cas NS1 retrovirus]	3e-14	
					PROTO-ONCOGENE C-CBL		
		Human mRNA for c-			human >gi 29731 (X57110) c-		
1327	X57110	cbl proto-oncogene	le-07	115855	cbl protein [Homo sapiens]	4e-19	
		Homo sapiens					
		(subclone 2_g12 from					
		BAC H94) DNA	4- 00	-MONTE-	<none></none>	<none></none>	
1328	AC001178	sequence Human interleukin-8	4e-08	<none></none>	SINOINES	CHOILE	
		receptor type B			1		
		(ILSRB) gene,					
		promoter and exons 1-		, ·			
1329	U11866	6	4e-08	<none></none>	<none></none>	<none></none>	
1327	0.1.000	Homo sapiens					
1		(subclone 2_e6 from			·		
		BAC H94) DNA			histone H1 II-1 (clone L95) -		
1330	AC001225	sequence	4e-08	478184	midge	6.5	
					HYPOTHETICAL 32.6 KD PROTEIN IN TRANSPOSON		
					TN4556 >gi 80758 pir JQ0428		
		Human modulator			hypothetical 32.6K protein -		
		recognition factor 2			Streptomyces fradiae transposon		
	1472027	(MRF-2) mRNA,	4e-08	141448	Tn4556	4.7	
1331	M73837	complete cds. Homo sapiens clone	46-08	141440	1114930		
		UWGC:y28gap from					
		6p21, complete			(AF000996) ubiquitous TPR		
		sequence (Homo			motif, Y isoform [Homo		
1332	AC006164	sapiens]	4e-08	2580578	sapiens]	1.2	
					T-CELL RECEPTOR BETA		
		Human mRNA for			CHAIN PRECURSOR	1	
1333	X01060	transferrin receptor	4e-08	135514	precursor (ANA 11) - rabbit	0.61	
	Ī .				INSULIN RECEPTOR-		
				•	RELATED PROTEIN PRECURSOR (IRR) (IR-		
					RELATED RECEPTOR)		
122.	V10607	H.sapiens INE2	4e-08	124909	>gi 186555 sapiens	0.14	
1334	Y10697	mRNA Rattus norvegicus	46-09	124707	>gijtoobbb sapiciis j	<u> </u>	
		myr 6 myosin heavy					
;		chain mRNA.			myosin I, high molecular weight		
1335	U60416	complete cds	4e-08	102189	- Acanthamoeba sp	3e-08	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					HYPOTHETICAL 55.2 KD		
					TRP-ASP REPEATS	1	
1 1		Drosophila			CONTAINING PROTEIN		
1 1		melanogaster putative			T10F2.4 IN CHROMOSOME	1	
		GTP-binding_			III protein; similar to G-Beta]	
		regulatory protein			repeat region (Trp-Asp	1	
		beta chain (GPB)			domains) of guanine nucleotide		
1336	U23804	mRNA, partial cds.	4e-08	2494916	binding protein	le-28	
		Escherichia coli K-12			(AL022325) tF27C3.1.1		
1 1		MG1655 section 103			(protein similar to C. elegans		
		of 400 of the			protein B0035.16) (isoform 1)		
1337	AE000213	complete genome	4e-08	3294172	[Homo sapiens]	2e-67	
					RHO-RELATED GTP-		
		Mus musculus mRNA	• •		BINDING PROTEIN RHOD		
		for RhoM, complete			(RHO-RELATED PROTEIN		
1338	D89821	cds	2e-08	3024539	HP1) (RHOHP1) sapiens]	Ie-04	
1 1		**					
		Human telomeric					
		repeat DNA-binding					
		protein (PIN2)		MONT.	NONE	<none></none>	
1339	U74382	mRNA, complete cds	1e-08	<none></none>	<none></none>	KNOINES	
		Homo sapiens (subclone H8 5_a10					
		from Pt 35 H5 C8)				ŀ	
1340	L35657	DNA sequence.	le-08	<none></none>	<none></none>	<none></none>	
1340	L33037	Human succinate	16-06	CHOILE	(AF060886) adenine	2.0	
		dehydrogenase			phosphoribosyltransferase		
1341	L21936	flavoprotein subunit	1e-08	3201678	[Leishmania tarentolae]	4.0	
1371		Homo sapiens gene		32010,0			
		for osteonidogen.			tritin - wheat		
1342	AB009777	promoter region	le-08	479388	>gi 391929 gnl PID d1003454	2.2	
		p. c.moter . eg.ro			GLUCOSE-6-PHOSPHATE		
181		Human heparin	1		ISOMERASE, CYTOSOLIC 2		
		cofactor II (HCF2)	ļ		(GPI) (PHOSPHOGLUCOSE		
 		gene, exons I through	I		ISOMERASE) (PGI) isomerase		
1343	M58600	5.	le-08	1730173	[Clarkia concinna]	1.9	
					GLUCOSE-6-PHOSPHATE		
		Human heparin	ļ		ISOMERASE, CYTOSOLIC 2		
		cofactor II (HCF2)	}		(GPI) (PHOSPHOGLUCOSE		
 		gene, exons I through			ISOMERASE) (PGI) isomerase		
1344	M58600	5	le-08	1730173	[Clarkia concinna]	1.7	
		Homo sapiens					
		(subclone 1_g2 from	}				
[PI H31) DNA	1		(L27428) reverse transcriptase		
1345	AC000980	sequence	le-08	439877	[Homo sapiens]	1.1	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	110210301						
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
	ACCESSION	DESCRIPTION			Ì		
\vdash		Human non-muscle					
		alpha-actinin mRNA.			(M76546) hydroxyproline-rich		
1346	U48734	complete cds	Ie-08	168237	protein [Helianthus annuus]	0.19	
	<u> </u>	Human leukocyte					
		adhesion receptor			(X92485) pval [Plasmodium		
1347	M76724	alpha subunit	1e-08	1177607	vivax]	0.19	
		6.11					
1		Gallus gallus homeodomain protein			ì	l	
		HOXD-3 mRNA.			(AF067942) No definition line		
1348	AF067959	complete cds	le-08	3165574	found [Caenorhabditis elegans]	0.15	
1346	A1 007939	Human DNA					
]		sequence from	٠.				
		cosmid U65A4.			İ	1	
		between markers					
		DXS366 and DXS87			(U93569) putative p150 [Homo	0.001	
1349	Z81014	on chromosome X *	1e-08	2072964	sapiens	0.001	
1		Human h-lys gene for					
1.250	V57.02	lysozyme (upstream	7e-09	<none></none>	<none></none>	<none></none>	
1350	X57103	region)	76-03	ZITOITES			
1			ŀ		BILE-SALT-ACTIVATED		
	Į				LIPASE PRECURSOR ESTER		
		Sambucus nigra	ĺ		LIPASE) (STEROL		
		ribosome inactivating			ESTERASE) (CHOLESTEROL		
	1	protein precursor			ESTERASE) salt-activated	0.22	
1351	AF012899	mRNA, complete cds	7e-09	231629	lipase [Homo sapiens] sapiens]	0.22	
		116			cytochrome-c oxidase (EC		
1		Aplysia californica prohormone			1.9.3.1) chain II precursor -		
		convertase (PC2)			Synechocystis sp. (PCC 6803)		
1352	L34741	mRNA, complete cds.	5e-09	322054	>gi 581739 sp.]	5.0	
1.232				·	THE TOTAL CONTROL OF THE TOTAL CONTROL OT THE TOTAL CONTROL OF THE TOTAL	· _	
					PHOTOSYSTEM II P680		
}					CHLOROPHYLL A		
	ł ·		1		APOPROTEIN (CP-47 PROTEIN)		
1	ĺ		į		>gi 7270S pir QJLV6A		
		**	1	ł	photosystem II chlorophyll a-		
		Homo sapiens type XV collagen			binding protein psbB - liverwort		
		(COLISAI) gene.			(Marchantia polymorpha)]	
1353	AF052959	exon 6	4e-09	131269	chloroplast >gi 11700	1.8	

	Nearest N	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	. 04,03(1						
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					POSSIBLE AGMATINASE		
					CAGMATINE		
					UREOHYDROLASE) (AUH)	}	
					(PROCLAVAMINIC ACID	1	
l					AMIDINO HYDROLASE)	1	
					>gi 1361423 pir \$57669	1	
					Proclavaminic acid amidino	1	
					hydrolase - Streptomyces	1	
!		Streptomyces		•	clavuligerus >gi 295171		
		clavuligerus (NRRL			Proclavaminic acid amidino	į į	
		3585) clavulanic acid			hydrolase [Streptomyces		
		biosynthesis protein			clavuligerus]	,	
		(cla) gene, complete			>gi 1586122 prf 2203286B		
		eds and clavaminate	٠.		proclavaminic acid amidino		
		synthase 2 (cs2) gene.			hydrolase [Streptomyces		
1354	L15470	partial eds.	4e-09	586028	clavuligerus	4e-13	
					GENERAL SECRETION		
l I		II			PATHWAY PROTEIN L		
		Human mRNA for			product [Klebsiella pneumoniae]		
1355	AB002302	KIAA0304 gene.	2e-09	131600	>gi 149311 (M32613) pulL	2.5	
1333	AB002302	complete cds	26-09	131000	>gij149511 (M52015) pai2	2.5	
		Homo sapiens				İ	
		retinaldchyde-binding					
		protein (CRALBP)			ļ		
1356	L34219	gene, complete cds.	le-09	<none></none>	<none></none>	<none></none>	
		Human mRNA for				i .	
		KIAA0304 gene.			(AB002302) KIAA0304 [Homo		
1357	AB002302	complete cds	le-09	2224549	sapiens]	5.0	
		**					
		Homo sapiens					
		HSPAIL gene for Heat shock protein 70					
[]	•	testis variant, 5'UTR.	-		(U58658) unknown (Homo	·	
1358	D85731	partial sequence	1e-09	1389766	[sapiens]	1.3	
1000	167797	partial sequence		2307700	[]		
		Homo sapiens natural				[
		resistance-associated			}		
		macrophage protein 2					
		(NRAMP2) gene.					
		exon 17, alternatively					
		spliced non-IRE			!!!! ALU CLASS F WARNING		
1359	AF064483	form, complete cds	8e-10	113671	ENTRY !!!!	0.72	

PCT/US00/18374

	Nearest N	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlustX vs. Non-Redundant Proteins)			
SEQ							
- 1	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
	7.0000.01.						
							
		Mus musculus alpha-			1		
		actinin-2 associated				1	
		LIM protein mRNA.			(AF053367) carboxyl terminal		
		alternatively spliced			LIM domain protein (Mus		
1360	AF002283	product, complete cds	6e-10	2996196	musculus]	4e-21	
		African green					
		monkey origin of			gene DMR-N9 protein - mouse		
1361	M26220	replication	5e-10	2143455	(fragment)	8.8	
		H.sapiens flow-sorted					
		chromosome 6		•	(1107574)		
		HindIII fragment,	4- 10-	2072077	(U93574) putative p150 [Homo	0.005	
1362	Z78006	SC6pA7F10	4e-10	2072977	sapiens] (U88183) similar to the	0.003	
					immunoglobulin superfamily,		
1 1		Homo sapiens			most similar to nerual cell		
		unknown protein			adhesion proteins		
1363	U82303	mRNA, partial cds	2e-10	1825711	[Caenorhabditis elegans]	0.031	
1303	002303	Drosophila	30.0				
		melanogaster			(AF079764) enhancer of		
l i	·	enhancer of			polycomb [Drosophila		
1364	AF079764	polycomb	2e-10	3757890	melanogaster]	le-10	
					(AC004520) similar to NFE2-		
					related transcription factors;		
1 1		Homo sapiens NRF1	1		similar to I48694		
		protein (NRF1)	. 1		(PID:g2137676) [Homo		
1365	L24123	mRNA.	2e-10	3004573	sapiens]	4e-53	
					NADH-UBIQUINONE		
i i			1		OXIDOREDUCTASE CHAIN		
			ł		4 >gi 1085185 pir S52968	1	
			Į į		NADH dehydrogenase chain 4 -		
			. 1		honeybee mitochondrion	}	
		Orangutan alpha-			(SGC4) >gi 552446 (L06178)		
		globin gene duplicate			NADH dehydrogenase subunit 4		
1366	M91454	region.	le-10	464239	[Apis mellifera ligustica]	6.0	
1500	17171737	House mouse;					
		Musculus domesticus					
		brain mRNA for					
		SAP102, complete	l		(L31961) phosphoprotein [Mus		
1367	D87117	cds	6e-11	473912	cookii]	2.2	
		Homo sapiens					
		(subclone 2_h9 from					
		P1 H39) DNA	1				
1368	AC001002	sequence	5e-11	<none></none>	<none></none>	<none></none>	

	Nearest	Neighbor (BlastN vs. C	ienbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC			JULIU 2011 R. /	Memest Mein	Newtest Neighbor (Brasta vs. Non-Redundant Proteins)		
19	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Homo sapiens		I			
		(subclone 2_h9 from					
1		P1 H39) DNA	ł		İ	1	
1369	AC001002	sequence	5e-11	<none></none>	<none></none>	<none></none>	
1		Homo sapiens					
1,270		KIAA0414 mRNA,					
1370	AB007874	partial cds	5e-11	<none></none>	<none></none>	<none></none>	
1		Homo sapiens	i i				
1	1	(subclone 2_h9 from				1	
1371	4.5001000	P1 H39) DNA	_			· .	
13/1	AC001002	sequence	5e-11	<none></none>	<none></none>	<none></none>	
		Homo sapiens		• •		•	
1		(subclone 2_h9 from				1	
1372	4001002	PI H39) DNA				1	
13/2	AC001002	sequence Homo sapiens	5e-11	<none></none>	<none></none>	<none></none>	
		(subclone 2_h9 from	Í	•		1	
		PI H39) DNA	J		1	1	
1373	AC001002	Sequence	5-11	NONE			
1313	AC001002	Homo sapiens	5e-11	<none></none>	<none></none>	<none></none>	
		(subclone 2_h9 from			1	1	
		PI H39) DNA			1 .		
1374	AC001002	sequence	5e-11	<none></none>	<none></none>	NONE	
		H.sapiens mRNA for	30-11	CHOILES	*NONE>	<none></none>	
		HERV-K long			gag polyprotein - human		
1375	Z21852	terminal repeat	5e-11	419481	endogenous virus S71	4.6	
					ondogenous views 571	7.0	
ŀ		Homo sapiens mRNA	ł			1	
		for KIAA0459	1		(AF051782) diaphanous I		
1376		protein, partial cds	5e-11	2947238	[Homo sapiens]	2.8	
- 1		House mouse:					
ı	ł	Musculus domesticus	j			i i	
j	i	brain mRNA for	1				
	1	SAP102, complete	·	•	(L31961) phosphoprotein [Mus	ľ I	
1377		cds	5e-11	473912	cookii]	1.8	
		Homo Sapiens DNA	į				
- 1		sequence between	j				
1270		two AML1 gene	1		!!!! ALU SUBFAMILY J		
1378	AJ131501	promoters, 6423 BP	5e-11	728831	WARNING ENTRY	0.20	
- 1	l.	Human andonosasus	l l			_ :-	
1	1	Human endogenous etroviral protease				}	
1379		• [5. 11	00650	retroviral proteinase-like protein	I I	
.5,7	171_70_0 1	nRNA, complete cds.	5e-11	88558	- human	0.002	

	Neurest	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	1							
1D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
<u> </u>					HYPOTHETICAL 55.2 KD			
1	1				TRP-ASP REPEATS			
	1	Drosophila			CONTAINING PROTEIN	1		
	ļ ·	melanogaster putative			T10F2.4 IN CHROMOSOME	1		
ĺ	ł	GTP-binding		'	III protein; similar to G-Beta			
Ĭ	İ	regulatory protein			repeat region (Trp-Asp	1		
1]	beta chain (GPB)			domains) of guanine nucleotide	ļ		
1380	U23804	mRNA. panial cds.	5e-11	2494916	binding protein	le-30		
		M.musculus troponin			(AF072889) transcription	1		
1381	Z22784	I gene.	3e-11	3892202	repressor brain factor 2	0.053		
	i	Homo sapiens						
		KIAA0420 mRNA,			1			
1382	AB007880	complete cds	2e-11	<none></none>	<none></none>	<none></none>		
		9 Homo sapiens BAX						
1202	A T00004	gene, exon 6, partial						
1383	AF020361	sequeпсе	2e-11	<none></none>	<none></none>	<none></none>		
		Homo sapiens DNA	1		GLYCOPROTEIN D			
1384	L35600	1 ' 1	20.11	117,1050	PRECURSOR gD [Bovine			
1364	E33000	sequence. Human organic anion	2e-11	1174952	herpesvirus []	0.25		
		transporting			(1050) 1) having and if a			
1385	U21943	polypeptide	2e-11	2738223	(U95011) brain-specific organic anion transporter	0-10		
-1000	921343	Homo sapiens		2130223	(AF053367) carboxyl terminal	9e-19		
		carboxyl terminal	i		LIM domain protein [Mus			
1386	U90878	LIM domain protein	2e-11	2996196	musculus]	4e-23		
		Human orphan			mascaras	46.23		
		nuclear receptor			i			
		(DAX1) gene,	1					
1387	U31929	complete cds	6e-12	<none></none>	<none></none>	<none></none>		
ſ		Human von	1					
- 1		Willebrand factor	1					
ł		gene, exon 1, 2, and			}			
		3, and three Alu						
1388	M25828	repetitive elements.	6e-12	<none></none>	<none></none>	<none></none>		
ļ			1					
J		Homo sapiens mRNA	- 1					
1,200		for KIAA0841						
1389	AB020648	protein, partial cds H.sapiens genes for	3e-12	<none></none>	<none></none>	<none></none>		
ļ			1					
		tumor necrosis factor (Tnfa) and	1	. [l		
1390			20.12	NONE				
1 390		ymphotoxine (Tnfb) Homo sapiens	2e-12	<none></none>	<none></none>	<none></none>		
- 1		kallistatin (PI4) gene.						
		exons 1-4, complete	}	}		1		
1391	1	eds	2e-12	<none></none>	MONES	NONE		
		-luman cosmid	20-12	CINOTAES	<none></none>	<none></none>		
1392		QLL2C9 from Xq28	2e-12	<none></none>	-NONE -	<none></none>		
	27.040	Service nomind-o	C-13	NONE>	<none></none>	CINOINE>		

	Nanros	Neighbor (BlastN vs. G	anhunk)	Nonco Najahha (Black V. a. Na - D. J J D.			
		Telgnoof (Blastin vs. O	enoank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		H.sapiens flow-sorted					
		chromosome 6					
	1	HindIII fragment,			hypothetical protein (L1H 3'	I	
1393	Z79007	SC6pA20E2	2e-12 .	106322	region) - human	1.5	
1		Human tyrosine					
1		kinase TXK (txk)			(M55524) ORF 4; putative	j	
1394	U34377	gene, exon 13.	le-12	151484	[Pseudomonas aeruginosa]	4.3	
1		Mus musculus apg-1	,				
I		gene for novel			ALKALINE PROTEINASE		
		member of heat shock			PRECURSOR (ALP) precursor	4	
		protein 110, promoter			fungus (Acremonium		
1395	D70845	region	le-12	113658	chrysogenum)	3.5	
		Human vascular	• • •		(AF069731) calmodulin-		
		endothelial growth			dependent protein kinase II beta		
1396	M63978	factor gene, exon 8.	1e-12	3982737	M isoform [Rattus norvegicus]	0.083	
i							
ľ		Homo sapiens			ĺ	1	
		lysosomal alpha-	Ĭ	*			
		mannosidase (manB)					
1397	U60266	mRNA, complete cds	8e-13	<none></none>	<none></none>	<none></none>	
		elegans cosmid	ŀ	,			
1			1		(AC002542) similar to C.		
		F11A10. complete	j		elegans F11A10.5; 80%		
		Sequence Caenorhabditis	1		similarity to Z68297		
1398		elegans)	7- 17	2202224	(PID:g1130619) [Homo		
1376	208291	Caenorhabditis	7e-13	2393734	sapiens]	5e-34	
		elegans cosmid	l		(AC002542) similar to C.		
		F11A10, complete		4	elegans F11A10.5; 80%		
]]	1	sequence	i		similarity to Z68297		
		[Caenorhabditis	i		(PID:g1130619) [Homo		
1399		elegans]	7e-13	2393734	sapiens]	3e-38	
		Human DNA	- '` ''	2070134	[aubicita]	٥٥٠٥	
	ļ	sequence from	l				
		cosmid L21F12B,	1				
		Huntington's Disease	I		į į		
		Region, chromosome	i	•			
	1	4p16.3, contains	1				
1400		EST.	6e-13	<none></none>	<none></none>	<none></none>	
		H.sapiens DAP-			(AB007143) ZIP-kinase [Mus		
1401	X76104	kinase mRNA	6e-13	2911154	musculus	0.007	
	i i	H.sapiens flow-sorted					
- 1	ļ	chromosome 6 TaqI				Į.	
	ı	ragment.	i		hypothetical protein (L1H 3'	l	
1402		SC6pA13G4	5e-13		region) - human	2e-06	
	I	Iomo sapiens DNA			(AC004136) hypothetical		
1403	L35600 s	equence.	3e-13		protein [Arabidopsis thaliana]	1.7	

	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neigh	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEO						T		
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
		Cloning vector						
		pKODT complete			(Z49966) F35C11.4			
1404	AF090452	sequence	2e-I3	3876730	[Caenorhabditis elegans]	7.8		
		Human gene for ATP						
		synthase alpha]		
1	1	subunit, complete cds			gag polyprotein - human	j		
1405	D28126	(exon 1 to 12)	2e-13	419481	endogenous virus S71	3.4		
		Homo sapiens				3.,		
		transcription factor			(AC004080) transcription factor			
1406	AF005219	HOXDI3	2e-13	2822166	HOXA13 [Homo sapiens]	5e-09		
		·						
1		Homo sapiens mRNA						
		for KIAA0758			(AB018301) KIAA0758 protein			
1407	AB018301	protein, partial cds	2e-13	3882237	[Homo sapiens]	le-23		
		Mus musculus apg-l						
1 1	1	gene for novel			ALKALINE PROTEINASE	;		
		member of heat shock			PRECURSOR (ALP) precursor			
		protein 110, promoter			fungus (Acremonium			
1408	D70845	region	le-13	113658	chrysogenum)	3.1		
		Homo sapiens				3.1		
		genomic DNA, 21q						
		region, clone:			(X15332) alpha-1 (III) collagen			
1409	AG000691	T171BG33	8e-14	930045	[Homo sapiens]	3e-04		
		Mouse mRNA for						
1 1		neuropsin, complete	ł		(AJ005641) serine protease			
1410	D30785	cds	8e-14	3559978	[Rattus rattus]	2e-12		
		Haemophilus	i			i		
		influenzae Rd section			(AL035064) queuine trna-			
1 1		25 of 163 of the	ľ		ribosyltransferase	ı		
1411	U32710	complete genome	8e-14	4106673	[Schizosaccharomyces pombe]	2e-38		
		I			hypothetical protein 2 - North			
i	}	Homo sapiens	İ		American opossum (fragment)			
		genomic DNA. 21q	1		>gi 897721 (Z48955) ORF-2,	- 1		
		region, clone:	ĺ		putative RT [Didelphis			
1412	AG000886	64EI1X19	7e-14	1363925	virginiana)	1.1		
			T					
	II	H.sapiens CpG DNA.	j					
		clone 71d11, forward	j		(AC002328) F20N2.6	· •		
1413	Z62664	read cpg71d11.ft1a.	7c-14	3953461	[Arabidopsis thaliana]	0.085		
			İ		· T			
		Homo sapiens mRNA	į		İ			
]	f	for KIAA0632			!!!! ALU CLASS C WARNING	1		
1414	AB014532	protein, partial cds	7e-14	113668	ENTRY !!!!	0.040		

[Negree	Neighbor (BlastN vs. G	enhank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		Treighbor (Blastity Vs. C	CHOARK)	Nearest Neigh	bor (BlastX. vs. Non-Redundant P	roteins)	
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
1		H.sapiens telomeric					
	i	DNA sequence, clone				1	
1		20PTEL004, read			(AB012223) ORF2 [Canis	ı	
1415	Z96478	20PTELOO004.seq	7 <u>e-1</u> 4	2981631	familiaris]	2e-04	
		Mus musculus					
1416	A 5100004	Pontin52 mRNA,					
1410	AF100694	complete cds	4e-14	<none></none>	<none></none>	<none></none>	
1		Sambucus nigra					
1		ribosome inactivating			1	i i	
1		protein precursor				i .	
1417	AF012899	mRNA, complete cds	4e-14	ANONIC	NO. TO		
 • • • • • • • • • • • • • • • • • • •	A1 012 699	mikiNA, complete cus	46-14	<none></none>	<none></none>	<none></none>	
1 .		Homo sapiens MLL					
		gene breakpoint	`			[
		cluster region, intron			!!!! ALU SUBFAMILY J	1	
1418	AF033349	l, partial sequence	3e-14	728831	WARNING ENTRY	9.3	
		Homo sapiens			Wilder Control	7.3	
1		(subclone 4_f6 from			1		
		P1 H54) DNA	j		extensin - almond >gi 20420	1	
1419	AC001526	sequence	3e-14	99861	(X65718) extensin	9.2	
		Sambucus nigra		•			
		ribosome inactivating					
1400	4 F012000	protein precursor			!!!! ALU SUBFAMILY \$B		
1420	AF012899	mRNA, complete cds	3e-14	728832	WARNING ENTRY	0.15	
]		ERUBINI AS PREGUESOR		
i		}	J		EPHRIN-A2 PRECURSOR		
ı			i		(EPH-RELATED RECEPTOR	ļ	
- 1		Mus musculus	1		TYROSINE KINASE LIGAND 6) (LERK-6) sapiens]		
- 1		Pontin52 mRNA.	i		>gi 2924761 (AC004258)		
1421		complete cds	2e-14	3913573		8.7	
				3713313	EPL6_HUMAN (Homo sapiens) EIB PROTEIN, SMALL [-	0.7	
j		,	ĺ		ANTIGEN (E1B 19K)	ŀ	
ļ	I		•		>gi 74142 pir Q1AD25 early	i	
I	1		l		EIB 21K protein II - human		
ł	į		ļ		adenovirus 5 >gi 58489		
- 1					(X02996) mRNA 5 first reading	ļ	
- 1	į:	Sambucus nigra	ł		frame [Human adenovirus type		
-	ļ.	ribosome inactivating			5] adenovirus type 5]	I	
J	1.	protein precursor	[>gi 209797 (J01969) 21 kD	j	
1422	AF012899 [nRNA, complete cds	9e-15	119040	protein	1.5	

	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		T	T	The section (Blase vs. 14011-reconsident Proteins)			
ED	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					transcription factor GATA-4,		
]		Sambucus nigra			retinoic acid-inducible - mouse		
1		ribosome inactivating		1	>gi 293345 (M98339) GATA-	ł	
1422	4 F012000	protein precursor			binding transcription factor		
1423	AF012899	mRNA, complete cds	8c-15	477102	[Mus musculus]	0.57	
		Canis familiaris LINE			-		
		I clement ORF2		}	harant sizet as i	ļ	
1424	AB012223	mRNA, complete cds	8e-15	92385	hypothetical protein - rat	0.000	
	11301222	Mus musculus	96-13	92303	(fragment)	0.003	
		Pontin52 mRNA.					
1425	AF100694	complete cds	3e-15	· · <none></none>	<none></none>	<none></none>	
		Human pHS1-2				SHORES	
		mRNA with ORF	• .	!	}		
		homologous to					
		membrane receptor			collagen alpha 3(IV) chain - sea		
1426	X12433	proteins	3e-15	422532	urchin	8.9	
		-		-	PROBABLE NUCLEAR		
ı			ì		HORMONE RECEPTOR		
ł					E02H1.7		
		Sambucus nigra			>gi 3875431 gnl PID e1344980		
- 1		ribosome inactivating protein precursor			(Z47075) similar to Zinc finger,		
1427		mRNA, complete cds	20.15	1252142	C4 type (two domains)		
1427	AI 012099	Human DNA	3e-15	1353143	[Caenorhabditis elegans]	5.0	
1	Î	sequence from	1				
I		cosmid L75B9.			1		
		Huntington's Disease					
		Region, chromosome			(L24521) transformation-related	i	
1428	Z69651	4p16.3	3e-15	403460	protein [Homo sapiens]	0.60	
					lg heavy chain precursor		
l		Sambucus nigra	i		(B/MT.4A.17.H5.A5) - bovine]	
· }		ribosome inactivating			>gi 440 (X62916) anti-	j	
		protein precursor	1		testosterone antibody [Bos	1	
1429		mRNA, complete cds	2e-15	108750	taurus]	1.1	
1.120		H.sapiens SMA3	_ [(X83299) SMA3 gene product		
1430	X83299	mRNA	2e-15	671530	[Homo sapiens]	0.32	
	j,	Human p300 protein					
J		nRNA, complete cds.]	1	
- 1		> :: gb 162297 162297				ł	
- 1		Sequence 1 from	ł		ELA-ASSOCIATED PROTEIN	1	
1431		patent US 5658784	2e-15	3024341	P300	0.019	
	0010//	Jateir US 3033/84	26-13	3024341	12000	0.019	

_	Near	est Neighbor (BlastN v	s. Genbank)	Nearest Ne	ighbor (BlastX vs. Non-Redundant	D
SE	- 1				Sales (Slass) 13. : 1011-Redundant	Proteins)
ID.	ACCESSI	ON DESCRIPTION	PVALUE	ACCESSION		P VALU
					HYPOTHETICAL 43.1 KD	
	J	}	i		PROTEIN CI6CIU.6 IN	
			1 '		CHROMOSOME III	
	1		1		>gi 3874384 gn PID e1344078	3
	1			1	EST EMBL C08256 comes	i i
	Ĭ	1	1	i	from this gene; cDNA EST	f
	Į		- 1	Į.	EMBL:C09941 comes from the	is
	1	Mouse MHC (Qa) (22		gene; cDNA EST yk340a10.3	
	1	k gene for class I			comes from this gene; cDNA	
1432	X16516	antigen, exons 4-8	le-15	2496897	EST yk340a10.5 comes from this gene [Ca	
	i			0.70037	uns gene (Ca	7e-08
1433		Chicken tensin			tensin - chicken >gi 212752	1
1433	M74165	mRNA, complete co	s. le-15	283920	(M74165) tensin	1
	Į	H.sapiens gene for			C-5 1057 (Clistii	2e-19
	ł	immunoglobulin		•		1
		kappa light chain	1 1			
434	X71893	variable region O4 and O5			i	1
	11/10/5	and O3	9e-16	<none></none>	<none></none>	<none></none>
- 1			.]		(Z84479) match: multiple	
			i 1		proteins; match: O00407	
- 1			1		Q12829 P22127 P36861	i 1
		Ì	1 1		Q40219; match: P70550	1 1
- 1		1	1		Q41022 P22125 Q08155	1 1
			1 1		P35286; match: P51148 P51147	}
		Human Rar protein	1 1		P35293 P36861 P35289; match: P35284 Q40217 P51152	i
35	U05227	mRNA, complete cds.	9e-16	3036779	P51157 P51158: match: Q41022	
- 1					131137 131138: match: Q41022	3e-06
-		Chicken erythrocyte	1 1			í
- 1		anion transport	1		(U23175) similar to anion	1
36	_M23404	protein (band3)	1		exchange protein	1
~ -	14123404	mRNA, complete cds.	9e-16	726403	[Caenorhabditis elegans]	Ie-28
		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	,		alpha-L-fucosidase (EC	-10-20
- 1		Rat mRNA for liver a-	İ		3.2.1.51) I precursor, tissue	1
- [1	L-Fucosidase (EC	i		human >gi 178409 (M29877)	1
37		3.2.1.51)	9e-16	(7:00	alpha-L-fucosidase precursor	
		.,,	96-10	67502	(EC 3.2.1.5) [Homo sapiens]	2e-29
	1.	Sambucus nigra	1	I		
	į:	ribosome inactivating				- 1
	Į.	protein precursor	ł		·	
8 . /	AF012899	mRNA, complete cds	8e-16	<none></none>	-NONE	
	ŢĮ.	Mus musculus brain			<none></none>	NONE>
1		mitochondrial carrier	1	į	1	
	l c	protein BMCP1	1	i		I
1				1		1
	(Bmcp1) mRNA, omplete cds	8e-16	10	(AF078544) brain mitochondrial	

	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		H.sapiens MN/CA9			!!!! ALU SUBFAMILY J		
1440	Z54349	GENE	5e-16	728831	WARNING ENTRY	0.002	
)		•			
		Mus musculus SH3 domain-containing			(M35536)		
1 1		adapter protein			(M35526) complement component CSD [Mus		
1441	AF077003	mRNA, complete cds	3e-16	309123	musculus]	3.1	
		M.musculus mRNA			glycoprotein - rat >gi 986943		
		for splicing factor			(L08134) glycoprotein [Rattus		
1442	X64587	U2AF (65 kD)	3e-16	2143767	norvegicus] norvegicus]	0.003	
		77					
		Homo sapiens mRNA for KIAA0661		•	(A BOLASSI) VIA A 0661	İ	
1443	AB014561	protein, complete cds	3e-16	3327136	(AB014561) KIAA0661 protein [Homo sapiens]	1e-20	
	.12014301	Human DNA	36-10	3327130	(Tonio sapiens)	10-20	
		sequence from					
		cosmid N120B6 on					
		chromosome 22					
]]	,	Contains ESTs.	•		·		
1444	Z73987	complete séquence	10.16	NONE	MONES	aNIONTT:	
1444	2/398/	[Homo sapiens] Homo sapiens ala	1e-16	<none></none>	<none></none>	<none></none>	
1445	M58318	gene.	le-16	<none></none>	<none></none>	<none></none>	
			i				
		Human small GTP binding protein Rab9			(280222) harrathariant anni-		
1446	Ŭ44103	mRNA, complete cds	1 c -16	1552584	(Z80233) hypothetical protein Rv0029	1.3	
	GIIIOS	mite vivi, complete cus	10 10	1332304		1.3	
		Homo sapiens mRNA	I				
l l	l l	for KIAA0661			(AB014561) KIAA0661 protein		
1447		protein, complete cds	9e-17	3327136	[Homo sapiens]	2e-20	
	1	Mus musculus Pontin52 mRNA.	1				
1448	1	complete cds	le-17	<none></none>	<none></none>	<none></none>	
. 7-75	1000,4	complete cas	10-17	21101122	CHOILE	×1,01,12	
ŀ	1	Mus musculus	ļ			ł	
J		ribosomal protein (Ke			1		
		3) gene, exons 1 to 5.			pupR protein - Pseudomonas		
1449	M76762	and complete cds.	le-17	1073048	putida >gi 525260	0.36	
		Human DNA.			LINE-I REVERSE		
.		replication enhancing			TRANSCRIPTASE		
1450		element (REE1)	4e-18	126295	HOMOLOG	0.78	
		Human mRNA for					
		hepatoma-derived					
		growth factor,	1				
1451	D16431	complete cds	4e-18	3242079	(AJ006984) proline-rich protein	0.018	

	Negreet	Naighbar (BlassN vs. C	anhants)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
050		Neighbor (BlastN vs. G	(ichoank)	New est Neighbor (DiastA vs. Non-Redundant Proteins)			
SEQ	1			}		ŀ	
B	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
							
ĺ					(Z73102) Similarity to B.subtilis		
ı			ľ		DNAJ protein	l	
1		Mus musculus heat			(SW:DNAJ_BACSU); cDNA	1	
1452	15000000	shock protein hsp40-3			EST yk437a1.5 comes from this		
1432	AF088983	mRNA, complete cds	4e-18	3873707	gene [Caenorhabditis elegans]	9e-25	
	ļ	Human methyl sterol					
İ		oxidase (ERG25)					
1453	U60205	mRNA, complete cds	3e-18	<none></none>	<none></none>	ANONE:	
		inita vit. complete das	30-10	CHOILD	CHOINES	<none></none>	
				٠,	G protein-coupled receptor 74 -		
1		Homo sapiens clone			equine herpesvirus 2 >gi 695246	i	
1		23899 mRNA			(U20824) G protein-coupled		
1454	AF038177	sequence	1e-18	1360775	receptor [Equine herpesvirus 2]	5.1	
		Homo sapiens mRNA	l		·		
1		for KIAA0661			(AB014561) KIAA0661 protein		
1455	AB014561	protein, complete cds	le-18	3327136	[Homo sapiens]	le-21	
1							
1		Homo sapiens mRNA					
1456	AB014561	for KIAA0661		2007106	(AB014561) KIAA0661 protein	1	
1430	AB014361	protein, complete cds	le-18	3327136	[Homo sapiens]	1e-22	
	`	Human tyrosine	· 1		1		
1 1		kinase TXK (txk)					
1457	U34374	gene, exons 9 and 10.	Ie-19	NONE	<none></none>	<none></none>	
		Homo sapiens				VIII OILES	
	1	hGAAI mRNA.	i		(AF102855) synaptic SAPAP-	ı	
1458		complete cds	le-19	4151809	interacting protein Synamon	0.19	
		Human mRNA for					
		KIAA0295 gene,	j		(AB002293) KIAA0295 [Homo	ľ	
1459	AB002293	partial cds .	1e-19	2224531 .	sapiens]	6e-17	
i 1	1		. 1	•	·		
} !		H.sapiens CpG DNA.	l				
1460		clone 16819, reverse			(Z82055) predicted using	I	
1400	Z59664	read cpg168f9.rt1a . Human modulator	5e-20	3880251	Genefinder	6.5	
			i			ĺ	
		recognition factor 2 (MRF-2) mRNA,	1				
1461		complete cds.	5e-20	201717	modulator recognition factor 2 -		
1401	1113031	complete cus.		284313	human factor 2 [Homo sapiens]	0.019	

-	Near	est Neighbor (BlastN vs.	Genbank)	Nearest Ne	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC	2				ignoof Blaste vs. Non-Redundant	Proteins)		
ID	ACCESSI	ON DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VAL		
1			 	+	DELTA-I-PYRROLINE-3-			
	ſ		ł		CARBOXYLATE			
	1	1	1		DEHYDROGENASE	1		
	1				PRECURSOR (P5C	1		
	į.	}	1	1	DEHYDROGENASE)	1		
	ı		1	j	>gi 1353248 sapiens]	ı		
	1		1		>gi 1353250 (U24267) pyrrolin	1		
	1	•	1	j	5-carboxylate dehydrogenase	9		
	}		1	j	[Homo sapiens]	1		
		Human pyrroline-5-	ļ		>gi 1589585 prf 2211355A	l		
1462	U24267	carboxylate	İ		Deltal-pyrroline-5-carboxylate	1		
. 102	024267	dehydrogenase	5e-20	2506350	dehydrogenase [Homo sapiens]	5- 04		
		Mus musculus myelin			(U13262) myelin gene	5e-04		
1463	U13262	gene expression factor			expression factor [Mus	ĺ		
	013202	Mus musculus myelin	4e-20	536926	musculus]	3e-07		
- 1		gene expression				30 07		
464	U13262	factor	4c-20	310/000	(AF061832) M4 protein			
T			46.70	3126878	deletion mutant [Homo sapiens]	le-08		
		H.sapiens CpG DNA.						
- 1		clone 48f10, forward			(7)007473			
465	Z61239	read cpg48f10.ft1a.	4e-20	1669601	(D88747) AR401 [Arabidopsis thaliana]			
- 1				1007001	(marrana)	8e-19		
- 1		Mus musculus						
- 1		junctional adhesion	1		1			
166	U89915	molecule (Jam)	ł		(U89915) junctional adhesion			
-	309913	mRNA, complete cds	1c-20	3462455	molecule [Mus musculus]	7e-11		
		Gallus gallus p52 pro-	1			,,,,,,,		
		apototic protein	- 1					
67	AF029071	mRNA, complete cds	7e-22	2500 :00	(AF029071) p52 pro-apototic			
Γ		Figure 4. Nucleotide	- /C-±±	2599492	protein [Gallus gallus]	le-1-5		
		sequence of the	j	·	. :			
		pKS36 1.797 kb	ı		(3/52/205)			
68	M25636	insert.	6e-22	1196398	(M21305) unknown protein			
- 1				1.70376	[Homo sapiens]	0.65		
		Homo sapiens mRNA	i	ł				
59 A		for KIAA0848	i	l.	(AB020725) KIAA0918 protein			
<u>~_</u>	B020655	protein, complete cds	6e-22	4240325	[Homo sapiens]			

	Nangast	Majakhar (BlassN vs. (Ganhank)	Name of the	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	1	Neighbor (BlastN vs. C	Jenoank)	Nearest Neigh	nbor (BlastX vs. Non-Redundant P	roteins)		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
					PROCOLLAGEN ALPHA	Ť –		
1					I(IV) CHAIN PRECURSOR			
	1		ļ		>gi 84917 pir A31893 collagen	1		
1				ŀ	alpha 1(IV) chain precursor -	1		
	ļ	1	i		fruit fly (Drosophila	1		
1	ĺ		I		melanogaster) melanogaster]			
1	[chorionic .	i		>gi 157078 (M96575) type [V	ł		
l	1	gonadotropin beta 1	Ī		collagen pro-collagen	ŀ		
1470	S80935	(CG beta 1) subunit	5e-22	115310	[Drosophila melanogaster]	0.027		
1		Homo sapiens						
1		microsatellite	i		!!!! ALU SUBFAMILY J			
1471	AF053066	D5S2926 sequence	2e-22	728831	WARNING ENTRY	3e-04		
		Danio rerio carbonic	٠.		CARBONIC ANHYDRASE			
1		anhydrase homolog]		(CARBONATE			
		CAH-Z mRNA.			DEHYDRATASE) >gi 2576335			
1472	U55177	complete cds	2e-22	3123190	(U55177) CAH-Z [Danio rerio]	5e-14		
ĺ	ł	Gallus gallus						
1473	15041050	ubiquitin specific			(AF016107) ubiquitin specific			
1473	AF064250	protease 66	2 e- 22	2736064	protease 41 [Gallus gallus] DRA PROTEIN (DOWN-	7e-37		
1		ļ			REGULATED IN ADENOMA)	İ		
					>gi 2135020 pir A47456 down-			
			1		regulated in adenoma (DRA) -			
			l		human >gi 291964 (L02785)			
					Nuclear localization signal at			
					AA 569-573, 576-580, 579-583;			
]]		Homo sapiens			acidic transer, activ. domain 620-			
		pendrin (PDS)			640,; homeobox motif 653-676			
1474	AF030880	mRNA, complete cds	2e-22	729367	[Homo sapiens]	4e-53		
		Mus musculus						
		Pontin52 mRNA,						
1475		complete cds	6e-23	<none></none>	<none></none>	<none></none>		
		Human mRNA for			Pm5 protein - human			
1476		pM5 protein	3e-23	107350	>gi 1335273 gnl PID e36241	le-04		
		Rattus norvegicus	1					
[]		PAD-R11 mRNA for	1		1			
		Peptidylarginine	j			1		
		deiminase type I,			1			
1477		complete cds	2e-23	<none></none>	<none></none>	<none></none>		
		Human h NAT allele	l l					
l		2-2 gene for	1					
		arylamine N-			(J04734) CDC6 protein			
1478		acetyltransferase	2e-23	171200	[Saccharomyces cerevisiae]	9.8		
	•	Human h NAT allele	i			1		
1		2-2 gene for arylamine N-	1			İ		
1479		arylamine N- acetyltransferase	20.22	171000	(J04734) CDC6 protein			
1:+/7	D109/1	accivitransierase	2e-23	171200	[Saccharomyces cerevisiae]	8.3		

		Near	est Neighbor (BlastN v	s. Genbank)	Nearest Ne	ighbor (BlastX vs. Non-Redundant F	
	S	EQ				ignoof (Blasex vs. Non-Redundant F	roteins)
	ļ.	D ACCESSI		1	ACCESSION	DESCRIPTION	P VALU
	-		Homo sapiens ML				11 17.20
	1,4	80 AF02454	AF4 fusion protein			serine/proline-rich FEL protein,	
	1-1-7	80 AF02454	II mRNA, partial cds	2e-23	2136142	splice form 1 - human	
			<u></u>			· · · · · · · · · · · · · · · · · · ·	le-20
	14	81 1.2222	Human AF-4 mRN	-	1	(AF031404) MLL-AF4 fusion	1.
	117	81 L13773		2e-23	3063962	protein [Homo sapiens]	le-20
		1	Mus musculus				16-20
	143	32 AF10069	Pontin52 mRNA.				i
	· · ·	72 10009	4 complete cds Drosophila	8e-24	<none></none>	<none></none>	<none></none>
	1	1	•	,			N.O.V.
		l l	melanogaster Rga a		J	1	į
	148	3 U75467	Atu genes, complete		·	(U75467) Atu [Drosophila	·
	1	073407	cds	8e-24	1658503	melanogaster)	2e-37
		1	Human HepG2 parti cDNA, clone	all			
	148	4 D17076	hmd5a09m5				
		3.7070	umananama	7e-24	<none></none>	<none></none>	<none></none>
		1	j	· ·		FMRFAMIDE-RELATED	
	1					NEUROPEPTIDES	
	1	1	Mus musculus	1 1		PRECURSOR >gi 416208	
	1	j	Pontin52 mRNA,	1 1		(U03137) neuropeptide	j
	148:	AF100694		7. 24		precursor FMRFamide-related	ł
		1	complete cus	7e-24	1169643	peptide [Lymnaea stagnalis]	7e-10
	l	j	Human 28S	.] [(Z81054) predicted using	
	1486	M11167	ribosomal RNA gene	. 2e-24	2075 (0)	Genefinder; Similarity to UDP-	
			Mus musculus	. 26-24	3875481	glucoronosyltransferases	5.1
	İ	i	Pontin52 mRNA,	1 1		luan, as as	
	1487	AF100694	complete cds	2e-24	549173	USPI PROTEIN PRECURSOR	- 1
			Cloning vector		349173	>gi 169623	1.2
ı		l	pAP3neo DNA.	1 1		(7/5225) 7	j
ļ	1488	AB003468	complete sequence	2e-24	987050	(X65335) lacZ gene product	
İ			Human mRNA of trk		207030	[unidentified cloning vector]	0.058
Į			oncogene > ::	1		(M74509) [Human endogenous	
l			gb 196186 196186	1		retrovirus type C oncovirus	
l			Sequence 23 from	, ,		sequence.], gene product [Homo	1
L	1489	X03541	patent US 5734039	2e-24	325465	sapiens]	·
			Homo sapiens			Suprems 1	3e-04
			(subclone 2_g11 from				1
	1 400	• • • • •	P1 H43) DNA	1		reverse transcriptase related	- 1
	1490	L81652	sequence	2e-24	225047	protein (Homo sapiens)	4- 13
	ı		Drosophila			protein (Fromo Sapiens)	4e-12
	- 1	•	melanogaster	1		j	ĺ
			strawberry notch				
,	491		(sno) mRNA,	- 1		(U95760) Sno [Drosophila	1
1	-771	U95760	complete cds	2e-24	2078282	melanogaster)	2e-41
	- 1		Mus musculus	T		(AF004835) tyrocidine	-0.41
ı	492		Pontin52 mRNA,			synthetase 3 [Brevibacillus	
•		ALTOUO94	complete cds	8e-25	2623773	brevis	8.6
							- · - 1

-						
<u> </u>	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant Pr	- croine'
Si	EQ				Diasar vs. Hon-Redundant Pi	roteins)
1	D ACCESSIO	ON DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-						
14	93 AB00240		8e-25	2496822	HYPOTHETICAL 127.3 KD PROTEIN B0416.1 IN CHROMOSOME X >gi 746502 (U23516) B0416.1 gene product [Caenorhabditis elegans]	
149	94 K03002	Human mRNA from chromosome 15 gene with homology to MHC-HLA-SB-1 intron A.		1514614	(X92842) nuclear protein [Mus musculus]	9e-11
149	95 U61232	Human tubulin- folding cofactor E mRNA, complete cds		1465772	(U61232) cofactor E [Homo	le-13 2e-05
149	6 U10245	Arabidopsis thaliana Col-O putative RNA helicase A mRNA, complete cds.	5e-25	1353239	(U10245) putative RNA helicase A [Arabidopsis thaliana]	le-37
149	7 X89211	H.sapiens DNA for endogenous retroviral like element Homo sapiens	- 3e-25	2065210	(Y12713) Pro-Pol-dUTPase	5e-06
1498	L81652	(subclone 2_g11 from P1 H43) DNA sequence	3e-25	2072961	(U93568) putative p150 [Homo sapiens]	5e-16
1499	X82895	H.sapiens mRNA for DLG2	2e-25	2497511	MAGUK P55 SUBFAMILY MEMBER 2 (MPP2 PROTEIN) (DISCS, LARGE HOMOLOG 2)	le-34
1500	M36654	Mouse homeo box 2.6 (Hox-2.6) mRNA, complete cds. Mus musculus (clone	9e-26	3323169	(AE001255) T. pallidum predicted coding region TP0854	1.9
1501	L36315	pMLZ-1) zinc finger protein	9e-26	1806134	(Z67747) zinc finger protein [Mus musculus]	4e-05
1502		Homo sapiens mRNA for KIAA0738 protein. complete cds	9e-26		!!!! ALU SUBFAMILY J WARNING ENTRY	1e-07
1 <i>5</i> 03		Homo sapiens putative transcription factor CR53	9e-26		ZINC FINGER PROTEIN ZFP-	le-17

	Nearesi	Neighbor (BlastN vs. (Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant F	Proteinc
SEC					The state of the s	Totellis)
ID	ACCESSIO		P VALUE	ACCESSION	DESCRIPTION	P VALUE
-	-	Homo sapiens				
1		(subclone 2_e6 from	1			
1.50		BAC H94) DNA	1		(U91823) small S protein	
1504	AC001225	sequence	8e-26	2653713	[Hepatitis B virus]	4.3
		Mus musculus			cyteine-rich surface antigen 72,	
1,505		Pontin52 mRNA,	•		CRP72 - Giardia lamblia	i
1505	AF100694	complete cds	8e-26	283446	(fragment)	3.4
1,500					!!!! ALU SUBFAMILY SQ	
1506	X94912	H.sapiens Pr22 gene	3e-26	728837	WARNING ENTRY	4e-09
1	j	Mus musculus	<u> </u>			
1,507	1	Pontin52 mRNA,	ľ		1	1
1507	AF100694	complete cds	2e-26	<none></none>	<none></none>	<none></none>
1						
	ł	Human small GTP				1
1.500	*****	binding protein Rab9			(AB014512) KIAA0612 protein	1
1508	U44103	mRNA, complete cds	1e-26	3327038	[Homo sapiens]	8.7
1					(AC005990) Contains repeated	
1	[region with similarity to]
ł					gb U43627 extensin (atExt1)	j
			j	•	gene from Arabidopsis thaliana.	
		Mus musculus			ESTs gb Z34165 and gb Z18788	
1,500		Pontin52 mRNA,	- 1		come from this gene.	
1509	AF100694	complete cds	9e-27	4056454	[Arabidopsis thaliana]	0.14
		Homo sapiens	- 1		LINE-I REVERSE	
		genomic DNA, 21q			TRANSCRIPTASE	
1510	1.0001010	region. clone:	I		HOMOLOG protein	
1310	AG001212	9H11N46	9e-27	126296	[Nyeticebus coucang]	0.012
			i			
		Mus musculus mucin	1			
1511	A E027121	glycoprotein MUC3			(U76551) mucin Muc3 [Rattus	
1311	AF027131	mRNA, partial cds	9e-27	2589172	norvegicus)	2e-14
		D	1			
1		Rattus norvegicus	ł			1
Ī		CTD-binding SR-like	ł		·	i
1512		protein r.49 mRNA.		•	(U49057) rA9 [Rattus	ľ
1312	U49057	complete cds	5e-27	1438534	norvegicus)	le-0-1
- 1	i.		1			
		Human, plasminogen	1		ĺ	1
1513		activator inhibitor-1	[1
1717		M.musculus partial	3e-27 .	<none></none>	<none></none>	<none></none>
- 1		-	1			
1211	1	cochlear mRNA			(Z78160) unknown [Mus	
.217	Z78160 (clone 28D2)	3e-27	1490362	musculus]	2e-05
			1			
- 1	I,	Janian C-C DM	1		(AB004538) LIPOIC ACID	i
ł		Lisapiens CpG DNA.		1	SYNTHETASE	· !
1515		lone 99b4, reverse			PRECURSOR(LIP-SYN)	
.515	204210 [F	ead cpg99b4.rtla	3e-27	2257538	[Schizosaccharomyces pombe]	1 c -06

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
		Homo sapiens				1	
		(subclone H8 6_h6				 	
	•	from P1 35 H5 C8)					
1516	L35659	DNA sequence.	le-27	<none></none>	<none></none>	<none></none>	
		Mus musculus				KINDINE	
		Pontin52 mRNA.			(U72686) odorant receptor 4	1	
1517	AF100694	complete cds	1e-27	1644471	[Danio rerio]	7.5	
		Mus musculus			(AF003534) hypothetical		
		Pontin52 mRNA,			protein 004L [Chilo iridescent		
1518	AF100694	complete cds	1e-27	2738388	virus]	6.7	
		Homo sapiens gene		•	(AL032636) Y40B1B.3	·	
1519	AB009271	for BCNT, partial cds	1e-27	3880909	[Caenorhabditis elegans]	4.6	
		Mus musculus					
		Pontin52 mRNA,			spermatophorin Sp23 - yellow		
1520	AF100694	complete cds	1e-27	2133579	mealworm molitor]	0.85	
		Mus musculus					
, , , , ,	. 5. 00.0	Pontin52 mRNA,		151005	ENDOGLUCANASE A		
1521	AF100694	complete cds	le-27	121805	PRECURSOR	0.58	
		Mus musculus Pontin52 mRNA.			(15035303)		
1522	AF100694		10.27	3722000	(AF035323) survival motor	0.10	
1322	AF100694	complete cds Mus musculus	le-27	3722000	neuron protein [Bos taurus]	0.10	
		Pontin52 mRNA.	1		(AF074902) laminin alpha chain		
1523	AF100694	complete cds	le-27	3328188	[Caenorhabditis elegans]	0.083	
	110 1000 /	comprete cus		7723.00	(Caerioriaadatis diegans)	0.005	
		Homo sapiens IkB			(AF074382) IkB kinase gamma		
1524	AF074382	kinase gamma subunit	le-27	3641280	subunit [Homo sapiens]	0.041	
					(AC005990) Contains repeated		
1					region with similarity to		
ı			1		gb U43627 extensin (atExt1)		
- 1	ļ]	Ī		gene from Arabidopsis thaliana.		
		Mus musculus	ĺ		ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA.	l		come from this gene.	· •	
1525		complete cds	le-27	4056454	[Arabidopsis thaliana]	6e-04	
		Homo sapiens	İ				
	1	(subclone 2_e10 from			2		
		P1 H49) DNA	1	20.50 :=	reverse transcriptase related		
1526	L78778	sequence	1e-27	225047	protein [Homo sapiens]	2e-09	
		U sing finger	f		ĺ	1	
-		Human zinc finger protein basonuclin	J		(U59694) zinc finger protein	Ī	
1527		mRNA, complete cds.	le-27	1488275	basonuclin [Homo sapiens]	9e-22	
/		mixiva, complete cus.	10-27	1400213	pasonucini (rionto sapiens)	20-22	
	į.	Human ribosomal			(AB004538) LIPOIC ACID		
- 1		protein L9 gene. 5'	j		SYNTHETASE		
		region and complete	1		PRECURSOR(LIP-SYN)		
1528		eds.	4e-28	2257538	[Schizosaccharomyces pombe]	2e-04	
1	<u> </u>				Hormeogacemaromy ces pomocj		

Į.	Nearest	Neighbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEO				realest reigi	1001 (Blasez Vs. 140n-Redundant Pr	oteins)	
ID D	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
	T		Ì			I VALUE	
					(Z46381) similar to lipoic acid		
İ			ł		synthase; cDNA EST yk283b6.3	1	
i		1	1		comes from this gene; cDNA	l	
1			i .		EST yk283b6.5 comes from this		
1	1		}		gene: cDNA EST yk472f5.3		
1	ł	H.sapiens CpG DNA,		ł	comes from this gene; cDNA		
		clone 99b4, reverse			EST yk472f5.5 comes from this	1	
1529	Z64210	read cpg99b4.rt1a.	4e-28	3 8 78570	gene; cDNA EST yk476e7.3	7e-11	
		Danio rerio carbonic			CARBONIC ANHYDRASE	76-11	
1 .		anhydrase homolog	ł		(CARBONATE		
		CAH-Z mRNA,	1	• 1	DEHYDRATASE) >gi 2576335		
1530	U55177	complete cds	4e-28	3123190	(U55177) CAH-Z [Danio rerio]	5e-21	
					(CSSTTT) CTATE (Build Tello)	20-21	
		Human mRNA for			ACYL-COA		
1 - 1		very-long-chain acyl-			DEHYDROGENASE, VERY-		
1 1		CoA dehydrogenase			LONG-CHAIN SPECIFIC		
		(VLCAD), complete			PRECURSOR (VLCAD)		
1531	D43682	cds	4e-28	1351839	>gi 930358 taurus]	3e-27	
1		Homo sapiens]		
		survival motor neuron			1		
		pseudogene. complete			!!!! ALU SUBFAMILY J		
1532	AF016591	sequence	3e-28	728831	WARNING ENTRY	3e-08	
		Mus musculus					
		Pontin52 mRNA,	i		!!!! ALU SUBFAMILY SB		
1533	AF100694	complete cds	2e-28	. 728832	WARNING ENTRY	2.5	
			1		DEHYDRIN DHN3		
			Í		>gi 100035 pir S18139 dehydrin		
		Mus musculus	i i		DHN3 - garden pea >gi 20709	Ī	
.53		Pontin52 mRNA,]		(X63063) pea dehydrin DHN3		
1534	AF100694	complete cds	2e-28	118588	[Pisum sativum]	0.004	
					FMRFAMIDE-RELATED		
					NEUROPEPTIDES		
	- 1	Mus musculus	ſ		PRECURSOR >gi 416208	j	
J			•		(U03137) neuropeptide		
1535		Pontin52 mRNA, complete cds	20.20	(1/0/13	precursor FMRFamide-related		
1333	AI 100094	complete cas	2e-28	1169643	peptide [Lymnaea stagnalis] (AC005990) Contains repeated	6e-04	
- 1	1		1		region with similarity to		
1	İ		- 1		-	- 1	
[1		gb U43627 extensin (atExt1) gene from Arabidopsis thaliana.	i	
- 1	<u>[</u> ;	Mus musculus			ESTs gb/Z34165 and gb/Z18788	- 1	
- 1		Pontin52 mRNA.					
1536		complete cds	2e-28		come from this gene.	0.05	
		ompiete eus		4030434	[Arabidopsis thaliana]	9e-05	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		. –					
B	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
igsquare					(AC005990) Contains repeated		
					region with similarity to		
			ļ !		gb U43627 extensin (atExt1)		
		•			gene from Arabidopsis thaliana.	F	
1 1	ı	Mus musculus			ESTs gb Z34165 and gbiZ18788	1 .	
		Pontin52 mRNA,		.000.00	come from this gene.		
1537	AF100694	complete cds	2e-28	4056454	[Arabidopsis thaliana] (AC005990) Contains repeated	2e-06	
					region with similarity to		
			1 [gb U43627 extensin (atExt1)		
					gene from Arabidopsis thaliana.	1	
		Mus musculus	ļ 1		ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA.	ļ 1		come from this gene.		
1538	AF100694	complete cds	2e-28	4056454	[Arabidopsis thaliana]	2e-09	
1338	AL 100094	complete cus	26-20	40,044	(AC005990) Contains repeated	20-09	
			i i		region with similarity to		
			!		gb U43627 extensin (atExt1)		
			i		gene from Arabidopsis thaliana.		
		Mus musculus			ESTs gb Z34165 and gb Z18788	1	
		Pontin52 mRNA.		•	come from this gene.		
1539	AF100694	complete cds	2e-28	4056454	[Arabidopsis thaliana]	1e-09	
					(AC005990) Contains repeated		
į 1					region with similarity to		
į I					gb U43627 extensin (atExt1)		
į i			1 1		gene from Arabidopsis thaliana.]	
		Mus musculus	 		ESTs gb Z34165 and gb Z18788]	
		Pontin52 mRNA,]		come from this gene.		
1540	AF100694	complete cds	2c-28	4056454	[Arabidopsis thaliana]	5e-10	
1					(AC005990) Contains repeated	ļ j	
					region with similarity to		
		•	1		gb U43627 extensin (atExt1)		
	İ				gene from Arabidopsis thaliana.		
. 1		Mus musculus			ESTs gb Z34165 and gb Z18788		
ا ـ ر ا		Pontin52 mRNA.	2. 20	1056.17.	come from this gene.	,, ,,	
1541	AF100694	complete cds	2e-28	4056454	[Arabidopsis thaliana] (AC002131) Strong similarity to	1e-11	
. 1		Mus musquitus	i		extensin-like protein gb[Z34465]		
		Mus musculus Pontin52 mRNA.			from Zea mays. [Arabidopsis		
1542			2e-28	3157926		8e-12	
1342		complete cds Mus musculus	26-20	3137920	thaliana)	00-1-	
		Pontin52 mRNA.			·		
1543		complete cds	le-28	<none></none>	<no:ne></no:ne>	<none></none>	
		Mus musculus		7	T		
		Pontin52 mRNA.					
1544		complete cds	1e-28	<none></none>	<none></none>	<none></none>	
		Mus musculus					
		Pontin52 mRNA.					
1545		complete cds	1e-28	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	1						
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
	1	Mus musculus	1	ACCESSION .		IF VALUE	
-		Pontin52 mRNA.					
1546	AF100694	[le-28	-NONT-	NONE		
1340	AF100694	complete cds Mus musculus	16-20	<none></none>	<none></none>	<none></none>	
		Pontin52 mRNA.			j		
1547	AF100694		1e-28	NONE	NONE		
1347	AF100094	complete cds Mus musculus	16-26	<none></none>	<none></none>	<none></none>	
1		Pontin52 mRNA.	1				
1548	AF100694	complete cds	le-28	<none></none>	ANIONITE >	ANONE	
1348	AP100094	Mus musculus	16-28	CNONES	<none></none>	<none></none>	
		Pontin52 mRNA.	1]		
1549	AF100694	complete cds	1e-28	··· <none></none>	NONE	NONT	
1349	AF100694	Mus musculus	16-26	KNONES	<none></none>	<none></none>	
	1	Pontin52 mRNA,] <u> </u>		}		
1550	AF100694	·	1e-28	<none></none>	NONTE	I NOTE	
1220	At 100094	complete cds Mus musculus	15-20	SNUMES	<none></none>	<none></none>	
		Pontin52 mRNA,					
1551	AF100694	complete cds	1e-28	<none></none>	NONE	ANONTE:	
1331	AI-100094	Mus musculus	16-20	STACINES	<none></none>	<none></none>	
ŀ		Pontin52 mRNA.					
1552	AF100694		1e-28	<none></none>	-NONE-	NONT	
1332	Al 100094	complete cds Mus musculus	16-20	KNONES	<none></none>	<none></none>	
		Pontin52 mRNA.				1	
1553	AF100694	complete cds	1e-28	<none></none>	<none></none>	<none></none>	
1333		Mus musculus	16-26	CHOILES	- KNONES	KINONES	
		Pontin52 mRNA.				1	
1554		complete cds	le-28	<none></none>	<none></none>	<none></none>	
	712 (000)4	Mus musculus	.0.50	41011E2	CITORIES	CIVOINE?	
i i		Pontin52 mRNA,	ſ		•		
1555	1	complete cds	1e-28	<none></none>	<none></none>	<none></none>	
		Mus musculus		42.101.125	11.01.22	1110112	
ł		Pontin52 mRNA.	1]	
1556		complete cds	1e-28	<none></none>	<none></none>	<none></none>	
		Mus musculus				1	
		Pontin52 mRNA,	j	Ì			
1557		complete cds	1e-28	<none></none>	<none></none>	<none></none>	
		Mus musculus					
1	1	Pontin52 mRNA.				l i	
1558		complete cds	le-28	<none></none>	<none></none>	<none></none>	
		Mus musculus					
		Pontin52 mRNA,	1	-			
1559		complete cds	le-28	<none></none>	<none></none>	<none></none>	
		Mus musculus					
		Pontin52 mRNA.					
1560	AF100694	complete cds	1e-28	<none></none>	<none></none>	<none></none>	
		Mus musculus					
1	Į1	Pontin52 mRNA.			٠.	1	
1561	AF100694	complete cds	le-28	<none></none>	<none></none>	<none></none>	

	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEO			T		Sor (Siasar 13) Frontieredunidant Pi	O(CIIIS)	
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION		
	I	Mus musculus	I VALUE	ACCESSION	DESCRIPTION	PVALUE	
		Pontin52 mRNA.				ļ	
1562	AF100694	complete cds	le-28	<none></none>	ANIONIE-	NONE	
1302	74 100034	Mus musculus	16-20	KNUNES	<none></none>	<none></none>	
]		Pontin52 mRNA.			ļ		
1563	AF100694	complete cds	le-28	<none></none>	<none></none>	<none></none>	
	12 100071	Mus musculus	.0 20	THOMES	CHOILES	SAONES	
		Pontin52 mRNA.					
1564	AF100694	complete cds	le-28	<none></none>	<none></none>	<none></none>	
		Mus musculus		47,07,125	THOME	CHOINES	
1		Pontin52 mRNA.	0_0				
.1565	AF100694	complete cds	le-28	. <none></none>	<none></none>	<none></none>	
		Human simple repeat	٠.		İ		
1566	M87708	polymorphism.	le-28	<none></none>	<none></none>	<none></none>	
		Mus musculus					
		Pontin52 mRNA,				1	
1567	AF100694	complete cds	le-28	<none></none>	<none></none>	<none></none>	
1					B; cDNA EST yk450d8.5 comes		
					from this gene; cDNA EST		
					yk249a6.5 comes from this		
					gene; cDNA EST yk219a2.5		
					comes from this gene; cDNA		
			- 1		EST yk355e4.5 comes from this		
ı	·				gene; cDNA EST yk224f4.5		
!					comes fr	Ī	
I					>gi 3924881 gn1 PID e1354569		
ſ		1	Í		from this gene; cDNA EST		
Į.	,		ĺ		yk249a6.5 comes from this		
1			1		gene; cDNA EST yk219a2.5	i	
1			i	•	comes from this gene; cDNA		
J		Mus musculus			EST yk355e4.5 comes from this		
		Pontin52 mRNA,			gene; cDNA/EST yk224f4.5		
1568	AF100694	complete cds	1e-28	3924779	comes from	3.0	
	[1		FMRFAMIDE-RELATED		
	- 1		ł		NEUROPEPTIDES		
		,	i		PRECURSOR >gi 416208	ŀ	
ŀ		Mus musculus	i		(U03137) neuropeptide	J	
ادحما		Pontin52 mRNA,			precursor FMRFamide-related		
1569	AF100694	complete cds	1e-28	1169643	peptide [Lymnaea stagnalis]	0.66	

	Nearest	Neighbor (Blast.N vs. C	ienbank)	Nearest Neigh	hbor (BlastX vs. Non-Redundant Pr	oteins)
SEQ						
D.	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE
			i		(-0.00000) summar to tammin	
					B, cDNA EST yk+50d8.5 comes	
					from this gene; cDNA EST	
1			l		yk249a6.5 comes from this	
1	i			•	gene; cDNA EST yk219a2.5	ł
1					comes from this gene; cDNA	l
					EST yk355e4.5 comes from this	j
					gene; cDNA EST yk224f4.5	
					comes fr	
1 1		,			>gi 3924881 gnl PID e1354569	
]					from this gene; cDNA EST	•
		·		•	yk249a6.5 comes from this	
1 1					gene; cDNA EST yk219a2.5	
ļ					comes from this gene; cDNA	
		Mus musculus			EST yk355e4.5 comes from this	
1570	A.E.100404	Pontin52 mRNA,		200	gene; cDNA EST yk224f4.5	
1570	AF100694	complete cds Mus musculus	Ie-28	3924779	comes from	0.65
		Pontin52 mRNA,		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	0.22	
1571	AF100694		1. 20	2122570	spermatophorin Sp23 - yellow	0.40
13/1	AF100694	complete cds Mus musculus	1e-28	2133579	mealworm molitor	0.49
1		Pontin52 mRNA.			spermatophorin Sp23 - yellow	
1572	AF100694	complete cds	le-28	2133579	mealworm molitor	0.49
15/-	14 100054	Mus musculus	10-20	2133373	cyteine-rich surface antigen 72,	0.49
		Pontin52 mRNA,			CRP72 - Giardia lamblia	
1573	AF100694	complete cds	le-28	283446	(fragment)	0.45
					SPERMATOPHORIN SP23	
		Mus musculus			PRECURSOR mealworm	
		Pontin52 mRNA,			>gi 161725 (M92928) structural	
1574	AF100694	complete cds	le-28	2498937	protein	0.33
1		Mus musculus				
- 1		Pontin52 mRNA,	i		(U60315) MC107L [Molluscum]	
1575		complete cds	1e-28	1492050	contagiosum virus subtype []	0.18
ĺ		Mus musculus	i			
	1	Pontin52 mRNA,		A.ac	spermatophorin Sp23 - yellow	
1576	AF100694	complete cds	le-28	2133579	mealworm molitor] DEHYDRIN DHN3	0.088
			1		>gi 100035 pir S18139 dehydrin	
- 1		Mus musculus	j		DHN3 - garden pea >gi 20709	
ſ		Pontin52 mRNA.	l		(X63063) pea dehydrin DHN3	
1577		complete cds	le-28	118588	[Pisum sativum]	0.018
	. 12 100094	complete cus	10-20	110300	DEHYDRIN DHN3	0.010
- 1		1			>gi 100035 pir S18139 dehydrin	
- 1		Mus musculus	1		DHN3 - garden pea >gi 20709	
- 1		Pontin52 mRNA.	I		(X63063) pea dehydrin DHN3	
1578	1	complete cds	1e-28	118588	[Pisum sativum]	0.016



	Nea	rest Neighbor (BlastN)	vs. Genbank)	Nearest Nai	ghbor (BlastX vs. Non-Redundant P	 _
SE	EQ.			Tourest Ive	Non-Redundant P	roteins)
n	D ACCESS	ION DESCRIPTION	N P VALUE	ACCESSION		P VALUE
-					DEHYDRIN DHN3	
		Mus =useulus			>gi 100035 pir S18139 dehydrir	1
		Mus musculus Pontin52 mRNA,		1	DHN3 - garden pea >gi 20709	
15	79 AF1006				(X63063) pea dehydrin DHN3]
1	7 74 1000	94 complete cds	1e-28	118588	[Pisum sativum]	0.012
	1		Ī	ł	(AC005990) Contains repeated	
1	1	ł	1	l	region with similarity to	l i
1	}				gb U43627 extensin (atExt1)	1
1	i i	Mus musculus		1	gene from Arabidopsis thaliana.	i ,
-	1	Pontin52 mRNA,	- 1	į	ESTs gb Z34165 and gb Z18788	1
158	0 AF10069		le-28	4056454	come from this gene.	1 1
			10-28	. 4030434	[Arabidopsis thaliana] DEHYDRIN DHN3	0.010
1	1		.			
		Mus musculus			>gi 100035 pir S18139 dehydrin	
1	ł	Pontin52 mRNA,			DHN3 - garden pea >gi 20709	
158	1 AF10069		1e-28	118588	(X63063) pea dehydrin DHN3	
				110300	[Pisum sativum] FMRFAMIDE-RELATED	0.002
	j		1		NEUROPEPTIDES	1
	į.		1 .		PRECURSOR >gi 416208	
1	ł	Mus musculus	-		(U03137) neuropeptide	1
		Pontin52 mRNA,			precursor FMRFamide-related	1
1582	AF100694	complete cds	1e-28	1169643	peptide [Lymnaea stagnalis]	0.002
1	İ				(AC005990) Contains repeated	0.002
	1	·	1 1		region with similarity to	
l	ł	İ	i i		gb U43627 extensin (atExt1)	- 1
]	ł		1		gene from Arabidopsis thaliana.	ł
	l	Mus musculus			ESTs gb Z34165 and gb Z18788	ſ
1583	AF100694	Pontin52 mRNA,	1 1	•	come from this gene.	j
.505	At 100094	complete cds	1e-28	4056454	[Arabidopsis thaliana]	0.002
	ļ		1 1		DEHYDRIN DHN3	
	l	Mus musculus	1 1		>gi 100035 pir S18139 dehydrin	j
		Pontin52 mRNA,			DHN3 - garden pea >gi 20709	i
1584	AF100694	complete cds	1 12 20	110000	(X63063) pea dehydrin DHN3	
		tompiete eus	1c-28	118588	[Pisum sativum]	0.002
i		j	1		(AC005990) Contains repeated	
	ı		1		region with similarity to	1
1			1 1		gb U43627 extensin (atExt1)	.
j		Mus musculus	1 1		gene from Arabidopsis thaliana.	1
		Pontin52 mRNA,			ESTs gb/Z34165 and gb/Z18788	ļ
1585	AF100694	complete cds	le-28	4056454	come from this gene.	
			 		[Arabidopsis thaliana] DEHYDRIN DHN3	200.0
- 1			1		>gi 100035 pir S18139 dehydrin	1
- 1		Mus musculus	1 1		DHN3 - garden pea >gi 20709	
	_	Pontin52 mRNA,			(X63063) peu dehydrin DHN3	ļ
1586	AF100694	complete cds	1e-28		rm.	0001
					Is someth Surface (III)	0.001

	Nearest	Neighbor (BlastN vs.	Genbank)	Negrest Neig	Nearest Neighbor (BlastX vs. Non-Redundant Pro		
SEC				rearest recig	noor (Blasee vs. Non-Reddingant Pr	oteins)	
B	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
	T			1	(AC005990) Contains repeated	FVALUE	
			· · · · · · · · · · · · · · · · · · ·	 	region with similarity to	 	
		}	1		gb[U43627 extensin (atExt1)		
1	1			ļ	gene from Arabidopsis thaliana.	ĺ	
ł		Mus musculus	ł	1	ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA,	1	1	come from this gene.		
1587	AF100694	complete cds	1e-28	4056454	(Arabidopsis thaliana)	0.001	
		Total Pictor Cas	1020	4030434	(AC005990) Contains repeated	100.0	
1			1	ł	region with similarity to		
	Į.			ļ	gb U43627 extensin (atExt1)		
1	l		Í	Ī	gene from Arabidopsis thaliana.		
ļ	l	Mus musculus		[ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA.	1	· ·	come from this gene.	·	
1588	AF100694	complete cds	le-28	4056454			
	10003	Complete eds	10-20	4030434	[Arabidopsis thaliana] (AC005990) Contains repeated	6e-04	
1	ł	ļ		ļ	region with similarity to		
ľ	}	1		Ĭ	gb U43627 extensin (atExt1)		
	ļ		1		gene from Arabidopsis thaliana.	i	
1	İ	Mus musculus			ESTs gb Z34165 and gb Z18788		
ł		Pontin52 mRNA.			come from this gene.	1	
1589	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]		
			10-20	7030434	(AC005990) Contains repeated	5e-04	
1					region with similarity to	i	
					gb U43627 extensin (atExt1)	Į.	
i i				•	gene from Arabidopsis thaliana.		
1		Mus musculus			ESTs gb Z34165 and gb Z18788		
i i		Pontin52 mRNA,			come from this gene.	I	
1590	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	5e-04	
					DEHYDRIN DHN3	30 0.1	
		18	1		>gi 100035 pir \$18139 dehydrin		
		Mus musculus	1 1		DHN3 - garden pea >gi 20709	i	
!!		Pontin52 mRNA,	1 1		(X63063) pea dehydrin DHN3	Į	
1591	AF100694	complete cds	1c-28	118588	[Pisum sativum]	2e-04	
					(AC005990) Contains repeated		
	ŀ				region with similarity to	1	
	ļ		1 1		gb U43627 extensin (atExt1)		
ı			} !		gene from Arabidopsis thaliana.	i	
	- 1	Mus musculus			ESTs gb Z34165 and gb Z18788	ļ	
		Pontin52 mRNA,]		come from this gene.		
1592	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana]	2e-04	
l					(AC005990) Contains repeated		
J	1	ı	[[region with similarity to		
	ł				gb U43627 extensin (atExt1)		
I	ĺ				gene from Arabidopsis thaliana.		
ı		Mus musculus	1		ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA,			come from this gene.		
1593	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana]	5e-05	



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	Negrest	Neighbor (BlastN vs.	Canbank)			
SEC		Tresgliool (Blasse Vs.	Genoank)	Nearest Neig	hbor (BlastX vs. Non-Redundant Pr	oteins)
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
					(AC005990) Contains repeated	
f					region with similarity to	
1	1	1			gb U43627 extensin (atExt1)	1
	1		1		gene from Arabidopsis thaliana.	
i		Mus musculus			ESTs gb Z34165 and gb Z18788	İ
1		Pontin52 mRNA,	1		come from this gene.	
1594	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana] (AC005990) Contains repeated	5e-05
ł	Į				region with similarity to	
1	ł ·				gb U43627 extensin (atExt1)	
1	1	,			gene from Arabidopsis thaliana.	
İ		Mus musculus			ESTs gb Z34165 and gb Z18788	
		Pontin52 mRNA,	1		come from this gene.	
1595	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	le-05
					(AC005990) Contains repeated	10-03
	Ĭ		, ,		region with similarity to	ĺ
1	i		1	•	gb U43627 extensin (atExt1)	1
}					gene from Arabidopsis thaliana.	i
		Mus musculus			ESTs gb Z34165 and gb Z18788	i
1		Pontin52 mRNA,]		come from this gene.	ľ
1596	AF100694	complete cds	1e-28	40 56 454	[Arabidopsis thaliana]	le-05
İ			} <u> </u>		(AC003990) Contains repeated	
1			1		region with similarity to	i
1			l 1		gb U43627 extensin (atExt1)	- 1
			ł ł		gene from Arabidopsis thaliana.	1
		Mus musculus	! !		ESTs gb Z34165 and gb Z18788	- 1
1507		Pontin52 mRNA,	1 1		come from this gene.	
1597	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	9e-06
					(AC005990) Contains repeated	
					region with similarity to	ļ
1 1			}		gb U43627 extensin (atExt1)	J
l		Mus musculus	ı		gene from Arabidopsis thatiana.	
	j	Pontin52 mRNA.			ESTs gb Z34165 and gb Z18788	[
1598	AF100694		1. 20	1056151	come from this gene.	
1270	AL 100094	complete cds	1e-28	4056454	[Arabidopsis thaliana] (AC005990) Contains repeated	6e-06
ı			1		region with similarity to	J
I					gb U43627 extensin (atExt1)	j
I]				1 '	
- 1	1	Mus musculus	- 1		gene from Arabidopsis thatiana. ESTs gb Z34165 and gb Z18788	I
	i i	Pontin52 mRNA.	1		come from this gene.	f
1599	1	complete cds	1e-28	4056454	[Arabidopsis thaliana]	5e-06
				1030434	[criaoloopsis thahana]	26-00
i		Mus musculus	}		RNA-BINDING PROTEIN	ŀ
1	l l	Pontin52 mRNA,			FUS/TLS protein [human.	- 1
1600	AF100694	complete cds	1e-28	544357	Peptide, 526 aa] [Homo sapiens]	4e-06
				- -	F act and drift (trottio advicted)	



1.	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ΙD	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					(AC005990) Contains repeated		
					region with similarity to		
					gb U43627 extensin (atExt1)		
[gene from Arabidopsis thaliana.]	
•		Mus musculus			ESTs gb Z34165 and gb Z18788	l	
		Pontin52 mRNA,			come from this gene.		
1601	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	2e-06	
					(AC005990) Contains repeated	i	
					region with similarity to]	
	:		j		gb U43627 extensin (atExt1)	İ	
		Mus musculus			gene from Arabidopsis thaliana. ESTs gb Z34165 and gb Z18788	Ì	
		Pontin52 mRNA.			come from this gene.	•	
1602	AF100694	complete cds	le-28.	4056454	[Arabidopsis thaliana]	2e-06	
1002	A.F 100094	complete cus	16-20.	4030434	(AC005990) Contains repeated	26-00	
]		region with similarity to		
					gb U43627 extensin (atExt1)		
i i					gene from Arabidopsis thaliana.		
		Mus musculus			ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA,	İ		come from this gene.		
1603	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana]	9e-07	
					(AC005990) Contains repeated		
[region with similarity to		
					gb U43627 extensin (atExt1)	1	
					gene from Arabidopsis thaliana.		
		Mus musculus			ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA,	. 22	.05	come from this gene.	0 0-	
1604	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana] FMRFAMIDE-RELATED	8e-07	
					NEUROPEPTIDES		
			1		PRECURSOR >gi 416208		
 	İ	Mus musculus			(U03137) neuropeptide		
	. 1	Pontin52 mRNA.	Ì		precursor FMRFamide-related		
1605		complete cds	1c-28	1169643	peptide [Lymnaea stagnalis]	7e-07	
- 303	14 100074	complete cas			(AC005990) Contains repeated		
	ļ		Ì		region with similarity to		
					gb U43627 extensin (atExt1)		
			ļ		gene from Arabidopsis thaliana.		
	i	Mus musculus			ESTs gb Z34165 and gb Z18788		
	•	Pontin52 mRNA.			come from this gene.		
1606	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana]	6e-07	
I					(AC005990) Contains repeated	ł	
			l		region with similarity to		
]	ļ	1		gb U43627 extensin (atExt1)		
İ			ĺ		gene from Arabidopsis thaliana.		
ŀ	1	Mus musculus	Ì		ESTs gb[Z34165 and gb[Z18788]		
		Pontin52 mRNA,	1. 20	1057.55	come from this gene.	£	
1607	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana]	5e-0	



1 (4.4	Nearest 1	Neighbor (BlastN vs. C	ienbank)	Nearest Neigh	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ								
Ð	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
					(AC005990) Contains repeated			
					region with similarity to			
			1	l	gb U43627 extensin (atExt1)			
					gene from Arabidopsis thaliana.	6 1		
		Mus musculus	1		ESTs gb Z34165 and gb Z18788			
		Pontin52 mRNA.	1		come from this gene.			
1608	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	3e-07		
					(AC005990) Contains repeated			
					region with similarity to	1		
					gb U43627 extensin (atExt1)			
					gene from Arabidopsis thaliana.	1		
		Mus musculus			ESTs gb Z34165 and gb Z18788			
		Pontin52 mRNA,	!		come from this gene.			
1609	AF100694	complete cds	1e-28-	4056454	[Arabidopsis thaliana]	. le-07		
					(AC005990) Contains repeated			
					region with similarity to	1		
1					gb U43627 extensin (atExt1)			
1					gene from Arabidopsis thaliana.	,		
		Mus musculus			ESTs gb Z34165 and gb Z18788			
	. /	Pontin52 mRNA,			come from this gene.			
1610	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	1e-07		
1 1			ŀ		(AC005990) Contains repeated			
					region with similarity to			
1					gb U43627 extensin (atExt1)			
1 1]		gene from Arabidopsis thaliana.			
		Mus musculus			ESTs gb Z34165 and gb Z18788			
		Pontin52 mRNA.		10001	come from this gene.	7- 09		
1611	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana] (AC005990) Contains repeated	7e-08		
					region with similarity to			
l 1					gb U43627 extensin (atExt1)			
					gene from Arabidopsis thaliana.			
		Mus museulus			ESTs gb/Z34165 and gb/Z18788			
		Mus musculus			come from this gene.			
ا دری ا		Pontin52 mRNA,	10.70	4056454	[Arabidopsis thaliana]	2e-08		
1612	AF100694	complete cds	1e-28	4030434	(AC005990) Contains repeated	2E-00		
					region with similarity to			
[gb U43627 extensin (atExt1)	1		
					gene from Arabidopsis thaliana.			
j ł		Mus musculus			ESTs gb Z34165 and gb Z18788			
		Pontin52 mRNA.			come from this gene.			
1613	AF100694		le-28	4056454	[Arabidopsis thaliana]	6e-09		
1017	WT.1000A4	complete cds	10-10	4030434	[[Wranigobara trianginal	00-07		

	Nearest i	Veighbor (BlastN vs. C	ienbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ					Sec. 2 in the second of the se	J.C.11.3,	
D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
	recession	DESCRIPTION	FVALUE	ACCESSION	(AC005990) Contains repeated	PVALUE	
<u> </u>					region with similarity to	ļ	
1			1		gb[U43627 extensin (atExt1)		
1					gene from Arabidopsis thaliana.	1	
		Mus musculus			ESTs gb Z34165 and gb Z18788	1	
l i		Pontin52 mRNA.			come from this gene.		
1614	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	5e-09	
		- Complete GG			(AC005990) Contains repeated	30.07	
:]		region with similarity to		
					gb[U43627 extensin (atExt1)		
1 1	•	1			gene from Arabidopsis thaliana.		
		Mus musculus			ESTs gb Z34165 and gb Z18788	· ·	
		Pontin52 mRNA,]		come from this gene.		
1615	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	4e-09	
					(AC005990) Contains repeated		
					region with similarity to		
					gb U43627 extensin (atExt1)		
			-		gene from Arabidopsis thaliana.		
		Mus musculus			ESTs gb Z3+165 and gb Z18788		
1616	, E100404	Pontin52 mRNA,		******	come from this gene.		
1616	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana] [AC005990] Contains repeated	7e-10	
					region with similarity to		
					gb U43627 extensin (atExt1)		
					gene from Arabidopsis thaliana.		
		Mus musculus			ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA.	ł		come from this gene.		
1617	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	6e-10	
					(AC005990) Contains repeated		
ŀ			1		region with similarity to		
Í	1	ļ	1		gb U43627 extensin (atExt1)	1	
	ļ		ļ		gene from Arabidopsis thaliana.		
.		Mus musculus			ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA,	ł	•	come from this gene.		
1618	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana]	5e-10	
			1		(AC005990) Contains repeated		
1	1		·]		region with similarity to		
į	İ		ł		gb U43627 extensin (atExt1)		
	l,	M	İ		gene from Arabidopsis thaliana.		
- 1		Mus musculus	i		ESTs gb Z34165 and gb Z18788	Į.	
1619		Pontin52 mRNA,	10.20	1026151	come from this gene.	ا ۱۰۰۸	
1019	AF100694	complete cds	1e-28	4036454	[Arabidopsis thaliana]	4e-10	

	Nearest	Neighbor (BlastN vs. C	ienhank)	Neprest Naigh	nbor (BlastX vs. Non-Redundant Pr	
SEO		THE RESERVE OF THE SECTION OF THE SE	T T	14671CS! 146181	ibol (Blaster vs. Non-Redundant Pr	oteins)
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
					(AC005990) Contains repeated	
1					region with similarity to	
1					gb U43627 extensin (atExt1)	l
			į.		gene from Arabidopsis thaliana.	[
]		Mus musculus]		ESTs gb Z34165 and gb Z18788	
		Pontin52 mRNA.			come from this gene.	
1620	AF100694	complete cds	1c-28	4056454	[Arabidopsis thaliana]	2e-10
1			_		(AC005990) Contains repeated	
		<u> </u>	}		region with similarity to	
l			i	1	gb U43627 extensin (atExt1)	
i				1	gene from Arabidopsis thaliana.	
ļ.	1	Mus musculus			ESTs gb Z34165 and gb Z18788	
		Pontin52 mRNA,	}	·	come from this gene.	
1621	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	5c-11
Ĭ i			ļ		(AC005990) Contains repeated	
					region with similarity to	
					gb U43627 extensin (atExt1)	
					gene from Arabidopsis thaliana.	
		Mus musculus			ESTs gb Z34165 and gb Z18788	
		Pontin52 mRNA,			come from this gene.	
1622	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana]	2e-12
		Petromyzon marinus				
1623	AF032896	polyadenylate binding			polyadenylate binding protein II	
1023	AF032896	protein	1e-28	1082703	human DEHYDRIN DHN3	2e-27
			,		>gi 100035 pir S18139 dehydrin	
		Mus musculus			DHN3 - garden peå >gi 20709	
		Pontin52 mRNA.			(X63063) pea dehydrin DHN3	ļ
1624	AF100694	complete cds	9e-29	118588	[Pisum sativum]	0.013
	12 100051	Mus musculus	7, 2,	110300	[[1 isuni sativuni]	0.013
ľ		Pontin52 mRNA.	1		spermatophorin Sp23 - yellow	•
1625	_ 1	complete cds	9e-29	2133579	mealworm molitor!	6e-04
						00 04
•		•]	į		(Z81071) predicted using	. [
l	1	ļ]		Genefinder; Similarity to	
- 1			J		Human small nuclear	
l	l		i		ribonucleoprotein E cDNA EST	j
ĺ		Mus musculus	·		yk375g7.5 comes from this	
1		Pontin52 mRNA,			gene; cDNA EST yk435f5.3	1
1626	AF100694	complete cds	9e-29	3876465	comes from this gen	9e-06
1					(AC005990) Contains repeated	
	ŀ	ļ	1		region with similarity to	1
i	İ	İ	1	•	gb U43627 extensin (atExt1)	
			1		gene from Arabidopsis thaliana.	ļ
•		Mus musculus	j		ESTs gb Z34165 and gb Z18788	ļ
		Pontin52 mRNA,			come from this gene.	
1627	AF100694	complete cds	8e-29	4056454	[Arabidopsis thaliana]	2e-06

Co !

	Nearest	Neighbor (BlastN vs. G	enhank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEO	1. Carest	Totalion (Diastry vs. O	I	i vernezi ivelăl	Tool (Blastix vs. Non-Redundant Pr	oteins)	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					ADP-RIBOSYLATION	T	
1	·				FACTOR 3 truit tly (Drosophila		
i i					melanogaster) >gi 507234	i	
1		Mus musculus			(L25063) ADP ribosylation	l	
		Pontin52 mRNA.			factor 3 [Drosophila		
1628	AF100694	complete cds	4e-29	728883	melanogaster]	0.016	
1						1	
		Mus musculus			RNA-BINDING PROTEIN		
1629	4 F100004	Pontin52 mRNA,	4 00	54405=	FUS/TLS protein [human.	l <u> </u>	
1629	AF100694	complete cds	4e-29	544357	Peptide. 526 aa] [Homo sapiens]	2e-07	
		•			(AC005990) Contains repeated	. ·	
					region with similarity to gb U43627 extensin (atExt1)		
			• •		12 '		
l		Mus musculus			gene from Arabidopsis thaliana. ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA.					
1630	AF100694	complete cds	4e-29	4056454	come from this gene. [Arabidopsis thaliana]	1- 00	
1030	AI 100034	Complete cus	46-29	4030434	ACYL-COA	le-08	
					DEHYDROGENASE, VERY-		
		Human mRNA for			LONG-CHAIN SPECIFIC		
		very-long-chain acyl-			PRECURSOR (VLCAD)		
		CoA dehydrogenase			dehvdrogenase precursor - rat		
		(VLCAD), complete	- 1		Acyl-CoA dehydrogenase		
1631	D43682	cds	4e-29	1168287	[Rattus norvegicus]	6e-37	
		M.tuberculosis accBC			(Z95556) accD1		
1632	Y07660	gene	4e-29	2113935	[Mycobacterium tuberculosis]	3e-47	
		Human alpha-satellite					
1		DNA from clone	j		1		
1633	X55367	pTRA-2.	1e-29	<none></none>	<none></none>	<none></none>	
		Homo sapiens					
	,	(subclone l_fl from	1		j		
		P1 H54) DNA	1				
1634		sequence	le-29	<none></none>	<none></none>	<none></none>	
		{Alu repeats, clone]		[
i		52H10} [human,	ſ			Ī	
1425		colonic mucosa.	, , ,	70000	!!!! ALU SUBFAMILY J		
1635	S75940	Genomic, 943 nt]	1e-29	728831	WARNING ENTRY	1e-07	
İ	1,	Homo sapiens	J		IIII AI II SUBEANU VI	Ĭ	
1636	i i	PACE-1 gene. exon 13	1e-29	770071	!!!! ALU SUBFAMILY J	7- 00	
.030	PROGRACI.	FACE+ gene. exon 13	16-29	728831	WARNING ENTRY	2e-09	
- 1	[,	Mus musculus SI-13					
		domain-containing	İ			Į	
	1	adapter protein					
1637		•	5e-30	<none></none>	<none></none>	<none></none>	
1637		nRNA, complete cds	5e-30	<none></none>	<none></none>	<none< td=""></none<>	



	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					(AC005990) Contains repeated		
					region with similarity to		
1 1					gb U43627 extensin (atExt1)]	
					gene from Arabidopsis thaliana.		
		Mus musculus .			ESTs gb Z34165 and gb Z18788		
		Pontin52 mRNA.			come from this gene.		
1638	AF100694	complete cds	4e-30	4056454	[Arabidopsis thaliana]	3e-10	
					·		
i I		Xenopus laevis			POLYADENYLATE-		
		poly(A)-binding			BINDING PROTEIN		
		protein (ABP-EF)		. (1)	polyadenylate-binding protein -		
1639	M27072	mRNA, complete cds.	4c-30	1352709	African clawed frog laevis	5e-21	
		B.taurus mRNA for	٠.				
		bovine vacuolar	-		(AF039573) abscisic acid- and		
1640	X58386	ATPase subunit A	2e-30	2773154	stress-inducible protein	4.3	
		M.tuberculosis accBC			(Z95556) accD1		
1641	Y07660	gene	1ċ-30	2113935	[Mycobacterium tuberculosis]	4e-47	
		G					
		Sus scrofa mRNA for			(AF007561) delta 6-desaturase		
	4 703 60 40	hypothetical protein	45.71	4102021	[Borago officinalis]	7.4	
1642	AJ236940	(5': clone 7C4) Homo sapiens	4e-31	4102021	[Borago Officinaris]	7.4	
		calcium-dependent					
		chloride channel- l					
		(hCLCA1) mRNA.			(AB017156) gob-5 [Mus		
1643	AF039400	complete cds	2e-31	3721912	musculus]	7e-08	
10.5	10007400	complete eds		<u> </u>	BOMBYXIN B-2 HOMOLOG		
		Homo sapiens	1		PRECURSOR silkmoth		
		(subclone 5_d9 from			>gi 217385 gnl PID d1003528		
		PI HI9) DNA			(D13924) Samia bombyxin		
1644	L77036	sequence.	1e-31	461663	homolog B-2 [Samia cynthia]	1.1	
		H.sapiens mRNA for					
		macropain subunit			(X61971) macropain subunit	-	
1645	X61971	delta	1e-31	296734	delta [Homo sapiens]	3e-06	
		human mitochondrial	1		(AC005990) Contains repeated		
		trnas and partial	j		region with similarity to		
		proteins 4 & 5;	Į		gb U43627 extensin (atExt1)		
		histidyl-, seryl-,	į		gene from Arabidopsis thaliana.		
		leucyl-trna genes:	į		ESTs gb/Z34165 and gb/Z18788		
ارير		urf4 and urf5	6. 70	4056464	come from this gene.	0000	
1646	L00016	(partial).	5e-32	4056454	[Arabidopsis thaliana] (AC005990) Contains repeated	0.002	
					region with similarity to		
			1		gb[U43627 extensin (atExt1)		
		Human acidic	Ì		gene from Arabidopsis thaliana.		
		ribosomal	Ī		ESTs gb Z34165 and gb Z18788		
		phosphoprotein P2	j	•	come from this gene.	j	
1647	M17887	mRNA, complete cds.	5e-32	4056454	[Arabidopsis thaliana]	12-05	
1047	141 1 1 0 0 1	maxx. complete cas.	36-32	7030434	If wraninghara mananal	10.00	

SEQ ID ACCESSION DESCRIPTION P VALUE ACC	ESSION DESCRIPTION P VALUE
Human mitogen- responsive	ESSION DESCRIPTION P VALUE
responsive	
, , , , , , , , , , , , , , , , , , ,	1 I
phosphoprotein DOC-	j
	(4.500.4503)
2 mRNA, complete	(AC004683) putative
1659 U53446 cds. 6e-34 33	95443 ammonium transporter. 3' partial 4.7
	PROTEIN-TYROSINE
	PHOSPHATASE EPSILON
	PRECURSOR (R-PTP-
Homo sapiens serine	EPSILON) >gi 1439605
protease mRNA.	(U62387) protein tyrosine
	07226 phosphatase-e [Mus musculus] 3.2
Human mitogen-	
responsive	
phosphoprotein DOC-	I FRION
2 mRNA, complete	LEP100 protein precursor -
1661 U53446 cds. 2e-34 10	4757 chicken >gi 212254 gallus 1.6
Homo sapiens	
endogenous retroviral	
sequence ERV-L pol	
gene, clone ERV-L	(AJ233597) reverse
1662 AJ233632 Human6 2e-34 38	60513 transcriptase [Mus famulus] 4e-10
Homo sapiens full	(AC002521) putative Ser/Thr
length insert cDNA	protein kinase [Arabidopsis
1663 AF086310 clone ZD51F08 8e-35 29	47070 (thaliana) 2.3 40S RIBOSOMAL PROTEIN
	S2 (STRINGS OF PEARLS
	PROTEIN)
	ribosomal protein S2 - fruit fly
	(Drosophila melanogaster)
Human mRNA for	melanogaster] >gi 515972
	0652 (U01335) ribosomal protein S2 2e-10
Homo sapiens mRNA	
for KIAA0565	(AB011137) KIAA0565 protein
1665 AB011137 protein, complete cds 3e-35 30	3654 [Homo sapiens] 2c-16
Human protease M	(AF091247) potassium channel
1666 U62801 mRNA. complete cds 2e-35 39	29231 [Rattus norvegicus] 1.0
Homo sapiens serine	
protease (Omi)	(AF020760) serine protease
	38915 [Homo sapiens] 9e-14



	Nearest l	Neighbor (BlastN vs. G	enbank)	. Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Human DNA					
		sequence from					
		cosmid U235H3 on			(M22333) unknown protein		
1668	Z93943	chromosome X	8e-36	1196432	[Homo sapiens]	3e-10	
					DEHYDRIN DHIN3		
					>gi 100035 pir S18139 dehydrin		
l i					DHN3 - garden pea >gi 20709		
					(X63063) pea dehydrin DHN3		
1669	X06778	Rabbit 18S rRNA	7e-36	118588	[Pisum sativum]	0.011	
		II					
i i		Homo sapiens mRNA, chromosome					
		1 specific transcript		,	(AE001350) hypothetical		
1670	AB007962	KIAA0493	3e-36 [.]	3329243	protein [Chlamydia trachomatis]	3.1	
1070	AB007902	Human DNA	36-30	3327243	protein (Cinamydia daenonians)	3.1	
		sequence from	1				
		cosmid U65A4.	1				
		between markers					
		DXS366 and DXS87			HYPOTHETICAL PROTEIN	i	
1671	Z81014	on chromosome X *	3e-36	141103	ORF-1137 mouse	0.038	
		Human DNA					
		sequence from					
		cosmid U65A4.			ļ	j	
ł		between markers					
	_	DXS366 and DXS87			(M29325) ORFI [Mus	0.006	
1672	Z81014	on chromosome X *	3e-36	198651	musculus]	0.006	
		Human transporter				ļ	
. [protein (g17) mRNA.			(U49082) transporter protein	1	
1673	U49082	complete cds	3e-36	1840045	[Homo sapiens]	2e-15	
13,3	047002	Human transcription	- 30 30	20.00-5	(
		factor SPI mRNA. 3'	j		HF-1 regulatory element binding		
1674	J03133	end.	3e-36	477133	protein - rat	2e-31	
	•						
		Homo sapiens mRNA				İ	
] <u> </u>		for KIAA0465	ł		(AB007934) KIAA0465 protein		
1675	AB007934	protein, partial cds	1e-36	3413892	[Homo sapiens]	4e-37	
		Mouse Hox-2.5			homeotic protein Hox B9 -		
1676	M34857	mRNA.	9e-37	106296	human (fragment)	0.15	
'		Homo sapiens	į		.	1	
		(subclone H8 5_a10	1			İ	
1,,,,	1.35/55	from P1 35 H5 C8)	00.37	2072060	(U93568) p-10 [Homo sapiens]	3e-05	
1677	L35657	DNA sequence. H.sapiens	9e-37	2072960	[Liomo sapiens]	36-03	
		endogenous				-	
		retrovirus HERV-	i		(Y17833) env protein [Human		
1678	X80240	KC4 DNA	8e-37	4185944	endogenous retrovirus K	1e-!5	
10/0	₩ 005+0	NC+ Una	00-27	7100777	lauroSaurona rerro runa rel		



1	Nearest N	leighbor (BlastN vs. Ge	nbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant Pro	teins)
SEQ						
D D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
	ACCESSION	Human DNA				
		sequence from				
		cosmid U235H3 on	1		hypothetical protein (L1H 3'	
1679	Z93943	chromosome X	9e-38	106322	region) - human	4e-13
1072	2,3,43				HYPOTHETICAL ZINC	
					FINGER PROTEIN ZK686.4	
					IN CHROMOSOME III	
					>gi 630780 pir S-14909 ZK686.4	
					protein - Caenorhabditis elegans	1
					>gi 304346 (L17337) coded for	
					by C. elegans cDNAs	
		٠.		•	GenBank:M88869 and T01933;	
		H.sapiens mRNA for			putative [Caenorhabditis	3- 37
1680	X97303	Ptg-12 protein	4e-38 ⁻	466044	elegans	3e-37
		H.sapiens mRNA for	2 20	2224220	SOP2-LIKE PROTEIN	5e-06
1681	Y08999	Sop2p-like protein	3e-38	3334339	SOF2-LIKE PROTEIN	50.00
		TI GO DNA			(U53181) F36D4.2 gene	
	•	H.sapiens CpG DNA,			product [Caenorhabditis	
	G-2005	clone 74g6, forward	2e-38	1245686	clegans}	0.19
1682	Z62887	read cpg74g6.ft1a. Human endogenous	26-38	1243000	Cicgano	
1	Ì	retrovirus clone				
1	}	c5.11. HERV-H		•		
1	1	multiply spliced	i			
١.	ļ	subgenomic leader.	Ì			
l	İ	protease and integrase			(Z14310) tripartite fusion	·
		region mRNA, partial			transcript PLA2L [Human	
1683	U35032	cds	1e-38	59977	endogenous retrovirus)	1e-06
		Human mRNA for			(AC00254-1) Unknown gene	
1		KIAA0220 gene.	1		product splice form-2 [Homo	
1684	D86974	partial cds	1e-38	3337386	sapiens]	8c-11
		Human nonmuscle] .			
1	1	myosin heavy chain		ì	(A DO22023) Tommusele musein	
		(NMHC) mRNA. 3'		4115745	(AB022023) nonmuscle myosin	2e-11
1685	M31013	end.	1e-38	4115748	heavy chain B	
		TT 2/2				
		Homo sapiens Arp2/3				
		protein complex		İ		
		subunit p20-Arc		·		
1,000	A F000003	(ARC20) mRNA.	4e-39	<none></none>	<none></none>	<none></none>
1686	AF006087	D.melanogaster crn	40-39		(AL009171) 62D9.a	
1607	. V50274	mRNA	4e-39	2655888	[Drosophila melanogaster]	4e-42
1687	X58374	nactor	 		GTP-BINDING PROTEIN	
	1	Human DNA for			TC10 ras-like protein [Homo	
1688	D85815	rhoHP1, complete cds	le-39	134080	sapiens}	3e-26



	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neigh	ibor (BlastX vs. Non-Redundant Pr	oteins)
SEQ ID	ACCESSION	. DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
1689	U49057	Rattus norvegicus CTD-binding SR-like protein rA9 mRNA, complete cds	4e-40	1438534	(U49057) rA9 [Rattus norvegicus]	5e-05
1690	Y08999	H.sapiens mRNA for Sop2p-like protein	4e-40	3334339	SOP2-LIKE PROTEIN	9 c -08
1691	AB002293	Human mRNA for KIAA0295 gene, partial cds	4e-40	2224531	(AB002293) KIAA0295 [Homo sapiens]	Ie-30
		Homo sapiens full length insert cDNA			DOUBLE-STRANDED RNA- SPECIFIC EDITASE 1 (DSRNA ADENOSINE DEAMINASE) (RNA EDITING ENZYME 1) >gi 1707502 gn PID e254627 (X99227) double-stranded RNA- specific editase [Homo sapiens] editase 1 hRED1-L [Homo sapiens] >gi 2039300 (U76421) dsRNA adenosine deaminase	
1692	AF086222 AF044127	Homo sapiens peroxisomal short- chain alcohol dehydrogenase (SCAD-SRL) mRNA, complete cds	1e-40	2829669 4105190	DRADA2b [Homo sapiens] (AF044127) peroxisomal short-chain alcohol dehydrogenase	0.6 I 2e-06
1694		Mus musculus Sil mRNA, complete cds	le-40	88608	SIL protein - human >gi 338088 (M74558) SIL	6e-23
1695	• 1	Mus musculus Sil mRNA, complete cds	1e-40	88608	SIL protein - human >gi 338088 (M74558) SIL	6e-23
1 6 96		Mus musculus Sil mRNA, complete cds	1e-40	88608	SIL protein - human >gi 338088 (M74558) SIL	5e-23
1697		Mus musculus Sil mRNA. complete cds	1e-40	88608	SIL protein - human >gi 338088 (M74558) SIL	5e-23
1698		Homo sapiens mRNA for K1AA0742 protein, partial cds	Ie-40	3882205	(AB018285) KIAA0742 protein [Homo sapiens]	6 e-31

	Nearest	Neighbor (BlastN vs. (Genbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant Pr	oteins)
SEC	ACCESSION	DESCRIPTION	P VALUE	ACCESSION .	DESCRIPTION	P VALUE
<u> </u>	-	 			ATP-BINDING CASSETTE	
	1	M.musculus abc2	J	!	TRANSPORTER ABC1 -	
1699	X75927	mRNA	le-40	728773	human >gi 495257 (X75926)	
		Mad III	10.40	120713	abc1 [Mus musculus]	3e-37
		Homo sapiens clone	j		(AF068195) putative	
		23954 mRNA			glialblastoma cell differentiation	
1700	AF038200	sequence	5e-41	3211975	related protein [Homo sapiens]	5e-14
		Human estrogen sulforransferase	[]			
		(STE) gene, exon 8				
1701	U20521	and complete cds	4e-41	<none></none>	<none></none>	<none></none>
					1 110112	CHOILES
	ł	Homo sapiens			-	
	1	branched chain alpha- ketoacid				
	l	dehydrogenase kinase	ŀ	•	[3-METHYL-2-	
		precursor, mRNA,	i	•	OXOBUTANOATE DEHYDROGENASE	
		nuclear gene			(LIPOAMIDE)] KINASE	
		encoding			PRECURSOR alpha-ketoacid	
1		mitochondrial	l		dehydrogenase kinase precursor	
1702	AF026548	protein, complete eds	2e-41	3182923	[Homo sapiens]	2e-09
1703	Y07660	M.tuberculosis accBC gene	2e-41	,	HYPOTHETICAL 66.5 KD PROTEIN F02A9.5 IN CHROMOSOME III >gi 280542 pir S28313 hypothetical protein F02A9.5 - Caenorhabditis elegans Genefinder: similar to Propionyl- CoA carboxylase beta chain; cDNA EST EMBL:M89018 comes from this gene; cDNA EST EMBL:D28069 comes from this gene; cDNA EST EMBL:D28068 comes from this gene; cDNA EST	3e-38
		Homo sapiens		403647	gene, CDNA EST	3e-38
		genomic DNA, 21q	ļ			
		region, clone:	ļ		hypothetical protein (L1H 3'	1
1704	AG001237 9	9H11N46	le-41		region) - human	5e-09
1705	AB007934	Homo sapiens mRNA for KIAA0465 protein, partial cds	le-41		(AB007934) KIAA0465 protein [Homo sapiens]	3e-12
	I	Iomo sapiens clone				<u> </u>
1704		24711 mRNA			İ	1
1706	AF055029 s	equence	5e-42	3250681	(AL024486) putative protein	2.2

	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neigh	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ								
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
					III-			
					PHOSPHATIDYLINOSITOL-			
					4,5-BISPHOSPHATE	İ		
i					PHOSPHODIESTERASE	ŀ		
					DELTA I (PLC-DELTA-I)	1		
1					(PHOSPHOLIPASE C-DELTA-	-}		
					1) (PLC-III) >gi 163538			
1		O.cuniculus mRNA			(M20638) phospholipase C-III	i		
1707	Z49747	for phospholipase C	5e-42	130227	[Bos taurus]	5e-36		
		Human set gene,				1		
1708	M93651	complete cds.	2e-42	<none></none>	<none></none>	<none></none>		
				•		, '		
]]		Sus scrofa mRNA for						
	0.	hypothetical protein			(U79010) delta 6 desaturase			
1709	AJ236940	(5': clone 7C4)	2e-42	2062403	[Borago officinalis]	8.5		
		Human erythroid						
	102424	differentiation protein	2 12	1700106	INHIBIN BETA A CHAIN			
1710	J03634	mRNA Mus musculus mRNA	2e-42	1708436	PRECURSOR	2e-10		
1711	A 13333777	i i	6- 12	2401017	STRIATIN	2 22		
1/11	AJ223777	for striatin Homo sapiens	6e-43	2494917	>gi 1495773 gnl PID e254158	2e-32		
]		potassium channel			(AEOIGIU) VCNI V2 IB IIIo-o			
1712	AF016411	subunit KCNA3.1B	2 c- 43	2708514	(AF016411) KCNA3.1B [Homo sapiens]			
1/1-	A1010411	Homo sapiens	26-43	2700314	sapiens	3e-13		
		(subclone 2_f10 from			hypothetical protein 3 - rat			
1713	AC001443	BAC 2913	1e-43	111814	>gil56589	2e-06		
17.5	110001443	BAC 2717	10 73	111014	MAGUK P53 SUBFAMILY	20-00		
li			ı		MEMBER 2 (MPP2 PROTEIN)			
		H.sapiens mRNA for			(DISCS, LARGE HOMOLOG			
1714	X82895	DLG2	6e-44	2497511	2)	6e-52		
		Human BENE			(X57960) ribosomal protein L7			
1715	U17077	mRNA, partial cds.	3e-44	53912	[Mus musculus] >ei 55489	8e-30		
	i	Homo sapiens mRNA	. 1	•	· '			
1716	AJ222700	for TSC-22 protein	2e-44	<none></none>	<none></none>	<none></none>		
					INHIBIN BETA A CHAIN			
1	ļ	j	ł		PRECURSOR PROTEIN)			
			l		(EDF) >gi 87936 pir B24248			
	1		1		inhibin beta-A chain precursor -]		
l	j	1	ł		human >gi 1\$1947 (J03634)			
1	[}	j		erythroid differentiation protein			
Į	Ì				precursor [Homo sapiens]	j		
1		Human erythroid	i		sapiens!			
ı	1	differentiation protein	}		>gi 226850 prt] 1608260B			
1717	J03634	mRNA	2e-44	124279	inhibin beta.A (Homo sapiens)	0.73		

						_
		Neighbor (BlastN vs. C	Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant P	roteins)
SE	~ (
LID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		T		Î		1. TALOI
1.		Homo sapiens mRNA				+
1		for KIAA0618	ı		(S80864) cytochrome c-like	1
171	8 AB014518	protein, complete cds	7e-45	1911548 -	[polypeptide sapiens]	1.6
					polypopined suprems)	1.0
1		H.sapiens genomic			(U29380) similar to adenylate	1
171	X76808	DNA clone d2	7e-45	86820 L	cyclase [Caenorhabditis elegans	2e-09
		Homo sapiens mRNA				2005
1	ļ	for beta 2-	1 .		(U95995) RNA-dependent RNA	
		microglobulin,	ļ		polymerase [Cryptosporidium	
1720	AB021288	complete cds	2e-45	2465521	parvum]	0.15
		H.sapiens mRNA for		٠.		
		transcription factor	ĺ			l .
172	X63468	TFIIE alpha	8e-46	<none></none>	<none></none>	<none></none>
1	1	Homo sapiens D2-2				
1722	AF019226	mRNA, 3'UTR	7e-46	<none></none>	<none></none>	<none></none>
		Human mRNA for	·			
		KIAA0064 gene.			HYPOTHETICAL PROTEIN	
1723	D31764	complete cds	2e-46	3123050	KIAA0064	le-15
ł	ļ	Human MHC class II				
i]	HLA-DR-beta-psi				
l]	(DW4/DR4)	j			
1		pseudogene, exons	- 1			
1724		3,4, 5,6, clones cosII-		41050.5	(Y17834) gag protein (Human	
1724	R02774	3301 and cosII-801.	le-46	4185946	endogenous retrovirus K]	2e-14
			1		BRIDE OF SEVENLESS	
1		ļ]		PROTEIN PRECURSOR	
i]	1	ľ		>gi 1079166 pir A47550 bride	i
ļ]	<u>'</u>	1		of sevenless precursor - fruit fly	Į
1725	X92109	H.sapiens hcgIX gene	9e-47	2498185	(Drosophila virilis) >gi 290216 virilis]	
	1371107	rasapiens negar gene	- Je-47	2490103	NADH-UBIQUINONE	1.4
		H.sapiens	ľ		OXIDOREDUCTASE CHAIN]
		mitochondrial DNA,			4 >gi 86696 pir A00435 NADH	.]
1726		complete genome	8e-47	128753	dehydrogenase (ubiquinone)	4e-15
		Human tumor			dentification (delinate)	46-12
		necrosis factor				1
1727	M85145	receptor, 3' flank.	3e-47	<none></none>	<none></none>	<none></none>
		H.sapiens			3.101.122	SIVOIVE
1		ndogenous	[J
	ſ	etrovirus HERV-			(Y17833) env protein [Human	1
1728	X80240 I	KC4 DNA	3e-47	4185944	endogenous retrovirus K1	7e-18
	T					
1		I.sapiens CpG DNA.	1			1
		lone 87t9, forward	1		(AE001222) T. pallidum	ł
1729	Z63594 r	ead cpg87f9.ft1a.	1e-47	3322743	predicted coding region TP0454	2.4

19	Negrect	Neighbor (BlastN vs. C	irnhank)	Noger Mein	hos (Black) as N = B	
		reignbor (Blastit vs. C	Jenoank)	inearest neigh	nbor (BlastX vs. Non-Redundant P	roteins).
ID SEC	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		R.ramus mRNA for		1		
		vascular type-1				
1		angiotensin II		1	(U43629) integral membrane	}
1730	X62295	receptor	4e-48	1209756	protein [Beta vulgaris]	le-07
1		Human tumor				
1		necrosis factor	1	i		
1731	M85145	receptor, 3' flank.	3e-48	<none></none>	<none></none>	<none></none>
1	į.		İ	1		
	1	Homo sapiens mRNA	ł	ì		i .
1,777		for KIAA0905		l	(AB020712) KIAA0905 protein	1
1732	AB020712	protein, complete cds	4e-49	4240299	[Homo sapiens]	2e-20
		Hama caniona —DNIA				1 1
1	1	Homo sapiens mRNA for KIAA0905			(1 1
1733	AB020712	protein, complete cds	2- 10	10.10000	(AB020712) KIAA0905 protein) • 1
11,33	AB020712	R.rattus mRNA for	3e-49	4240299	[Homo sapiens]	2e-20
1		vascular type-1				1
		angiotensin II			(II 13620) integral	j
1734	X62295	receptor	le-49	1209756	(U43629) integral membrane	١
-	1102275	icospio.	10-49	1209730	protein (Beta vulgaris)	7e-12
ĺ	1	Homo sapiens mRNA				į į
1		for E1B-55kDa-			(AJ007509) E1B-55kDa-	
1735	AJ007509	associated protein	le-49	3319956	associated protein	4e-24
					HYPOTHETICAL ZINC	
1			}		FINGER PROTEIN ZK686.4	
1			1		IN CHROMOSOME III	,
			i		>gi 630780 pir S44909 ZK686.4	-
			•		protein - Caenorhabditis elegans	i
			ı		>gi 304346 (L17337) coded for	1
					by C. elegans cDNAs	1
	l		I		GenBank:M88869 and T01933;	1
1776	1	H.sapiens mRNA for			putative [Caenorhabditis	.
1736		Ptg-12 protein Homo sapiens	1e-49	466044	elegans]	8e-3!
		homolog of Nedd5	I			
		(hNedd5) mRNA,	1		1	
1737		complete cds	4e-50	<none></none>	Nove	2102
		Homo sapiens	46-30	KNONES	<none></none>	<u> <non< u=""></non<></u>
		polycystic kidney	I		(L43619) polycystic kidney	
	4	disease (PKD1) gene,	l		disease 1 protein [Homo	•
1738		exons 35-42	4e-50	903758	sapiens]	3e-/ .
				702.50	Jupicita)	<u> </u>
Ì		Homo sapiens clone	1		1	!
ł		22 mRNA, alternative	1			
- 1		splice variant alpha-1.	ł		(AF009426) clone 22 [Homo	
1739	AF009424 c	complete cds	4e-50	2271473	supiens]	5e

-	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		I Straightful (Blasta visit C	I I	ivearest ivergi	ibor (Biasex vs. Noit-Redundant P	roteins)	
ID ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
<u> </u>					monosaccharid transport protein		
		Homo sapiens			STP4 - Arabidopsis thaliana	T	
1	l	(subclone 8_c11 from			>gi 16524 (X66857) sugar	1	
	1	P1 H22) DNA			transport protein (Arabidopsis	1	
1740	L77040	sequence.	2e-50	99758	thaliana)	6.4	
		Homo sapiens					
1		(subclone H8 5_a10				ļ	
	Y 0.5.55	from P1 35 H5 C8)				ŀ	
1741	L35657	DNA sequence.	2c-50	2072960	(U93568) p40 [Homo sapiens]	2e-05	
		Hama and CTC7					
1742	1100746	Homo sapiens CTG7a		MONTE			
1742	U80745	mRNA, partial cds Bovine mRNA for	le-50	<none></none>	<none></none>	<none></none>	
1743	D84514	1 1	1- 50	2070627	(AF103728) structural		
1743	D64314	p97, partial cds	le-50	3978527	polyprotein [Sindbis virus] LYSOSOMAL PROTECTIVE	9.9	
			l		PROTEIN PRECURSOR		
			ł		(CATHEPSIN A)		
		Human protective	ļ		(CARBOXYPEPTIDASE C)		
		protein mRNA.	ı		human >gi 190283 (M22960)		
1744	M22960	complete cds.	1e-50	131081	protective protein precursor	le-12	
					muf1 protein - human	10-12	
		H.sapiens mRNA for			>gi 762953 (X86018) mufl		
1745	X36013	MUF1 protein	le-50	1082610	[Homo sapiens]	1e-21	
		Human transcription					
_		factor LSF-ID	- 1		transcription factor LSF - human		
1746	U03495	mRNA, complete cds.	7e-51	2136296	>gi 476099	le-21	
		Homo sapiens	·				
	. 50.40	HRIHFB2157	[(AB015344) HRIHFB2157	·	
1747	AB015344	mRNA, partial cds	5e-51	3970874	[Homo sapiens]	2c-35	
1748	1402220	Human zinc finger	4-51	200	MYC-ASSOCIATED ZINC		
1740	M93339	protein mRNA. Human zinc finger	4e-51	3024110	FINGER PROTEIN sapiens]	2e-06	
		protein zfp6 (ZF6)			(AC007697) E19517 1 III.mg	I	
1749		mRNA, partial eds	4e-51	2689441	(AC003682) F18547_1 [Homo sapiens]	2e-11	
	0,1303	institution partial cus		2007441	60S RIBOSOMAL PROTEIN	26-11	
ı	j	1	1		L13A (23 KD HIGHLY BASIC	1	
ı	ļ	1	- 1		PROTEIN)	1	
ł	ŀ	1	1		>gi 345897 pir S29539 basic	1	
	ŀ	H.sapiens mRNA for			protein, 23K - human >gi 23691	1	
		23 kD highly basic	l		(X56932) 23 kD highly basic	- 1	
1750		protein_	4e-51	730451	protein [Homo sapiens]	12-11	
		H.sapiens flow-sorted					
		chromosome 6	Í				
1		HindIll tragment.	1				
1751	Z79054 :	SC6pA21E11	2e-51	<none></none>	<none></none>	<none></none>	

11 11 11	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ				rearest reign	bol (Blaset vs. Non-Redundant Fi	Oterns)	
B	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Homo sapiens					
		BAF60b gene, partial					
1752	AF068245	sequence	5e-52	<none></none>	<none></none>	<none></none>	
	ŀ						
		Sus scrofa mRNA for			RIBONUCLEOPROTEIN		
1,763		hypothetical protein			RB97D ribonucleoprotein	1	
1753	AJ236932	(5'; clone 4B8)	5e-52	400927	[Drosophila melanogaster]	4.7	
		Mus musculus scaffold protein Pbp1					
1754	A E002 (0.2	homolog mRNA,		2127124	(AF003693) scaffold protein		
1/34	AF003693	complete cds	6e-53	2197106	Pbp1 homolog [Mus musculus] CALMODULIN	2e-54	
1 1					1		
	1	Human calmodulin	• .		>gi 102408 pir JC1309		
1755	M27319	mRNA, complete cds.	5e-53	115528	calmodulin - Stylonychia lemnae		
1133	1.12/317	maxiva, complete cus.		113328	(SGC5) >gi 161195	0.002	
		Mouse house-keeping protein mRNA,			house-keeping protein - mouse		
1756	M74555	complete cds.	5 e-53	284775	>gi 193871	5e-30	
1 1	į		j		phosphoenolpyruvate		
			- 1		carboxykinase (GTP) (EC		
		77 70	I		4.1.1.32) precursor,		
		H.sapiens mRNA for	· 1		mitochondrial - human		
1757	X92720	phosphoenolpyruvate carboxykinase	6.51	2125015	carboxykinase (GTP) [Homo		
1/3/		Homo sapiens torsinB	6e-54	2135915	sapiens]	6e-21	
i 1		(DQ1) mRNA, partial	1		(A B002 105) I A K 1= (II		
1758		cds	2e-54	2760121	(AB002405) LAK-4p [Homo	0.3-	
		Mus musculus	20-34	2700121	sapiens]	0.27	
	1	B6CBA Lisch7	l		(U49507) Lisch7 [Mus		
1759		mRNA, partial cds.	2e-54	1236083	musculus	3e-27	
		Human DNA		122 2005			
		sequence from			(Y14657) hydrophobin	ſ	
		cosmid 92M18,	ſ		[Pleurotus ostreatus]	İ	
j	į.	BRCA2 gene region	Ī		>gi 2982620 gn1 PID e1283986	ı	
		chromosome 13q12-	i i		(AJ225061) POH2 hydrophobin	į	
1760		13.	1e-55	2370371	[Pleurotus ostreatus]	2.0	
1	3	Human cytochrome c					
f		oxidase subunit VIa	i			ŀ	
		gene, exon 3 and	1		(AE000682) hypothetical		
1761	U83702	complete cds	8e-56	2982994	protein [Aquifex aeolicus]	7.0	
- 1	,	Jome seeine DN	-				
- 1		Homo sapiens mRNA for transducin (beta)	ļ		(3/12701)	l	
1762		ike I protein	7e-36	1	(Y12781) transducin (beta) like	7. 70	
., 02	11-101 1	ike i protein	16-20	3021409	I protein [Homo sapiens]	7e-39	

		Neighbor (BlastN vs. C	ienbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU	
ļ	 						
1763	AB020673	Homo sapiens mRNA for KIAA0866 protein, complete cds	8e-57	2104553	(AF001548) Myosin heavy chain (MHY11) (5'partial) [Homo sapiens]	4e-04	
1764	AJ236932	Sus scrofa mRNA for hypothetical protein (5': clone 4B8) Human dystrophin	3e-57	400927	RIBONUCLEOPROTEIN RB97D ribonucleoprotein [Drosophila melanogaster]	4.7	
1765	L06900	gene, intron 1 containing pseudo exon. H.sapiens	le-58	. 4185129	(AC005724) unknown protein [Arabidopsis thaliana] thaliana]	· 7.0	
1766	X93334	mitochondrial DNA, complete genome	9e-59	1492050	(U60315) MC107L [Molluscum contagiosum virus subtype 1]	0.17	
1767	AF064856	Rattus sp. 7acomp protein mRNA, complete cds	3e-59	3169626	(AF064856) 7acomp protein [Rattus sp.]	2e-31	
1768	_	Homo sapiens alphatubulin isoform 1 mRNA, complete cds	2e-59	32015	(X06956) alpha-tubulin [Homo sapiens]	4e-22	
1769	775	Homo sapiens mRNA for FUS-CHOP protein fusion	le-60	746557	(U23523) histidine-rich [Caenorhabditis elegans]	0.45	
1770	AF013988	Homo sapiens serine protease mRNA.	1e-60	2564316	(AB006622) No similarities to any reported proteins [Homo sapiens]	0.26	
1771	1	Mus musculus ymphocyte specific nelicase mRNA. complete cds	7e-61	2137490	lymphocyte specific helicase - mouse musculus]	3e-25	
772	X93334 c	I.sapiens nitochondrial DNA, omplete genome	4e-61	. 70656	ubiquitin / ribosomal protein \$27a - human extension protein, HUBCEP80 [human. Peptide, 156 aa] ubiquitin extention protein [Cavia porcellus]	9e-08	
7 73	D38255 c	Iomo sapiens mRNA or CAB1, complete ds	4e-61		gene MLN 64 protein - human	4e-23	
774	ly h	fus musculus mphocyte specific elicase mRNA. complete cds	8e-62		lymphocyte specific helicase - mouse musculus]	Se-26	

1,50	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Human lipocortin-V			Human Annexin V With Proline		
1775	M21731	mRNA, complete cds.	6e-62	3212603	Substitution By Thioproline	2e-20	
		Rattus norvegicus	1				
1		myotonic dystrophy					
		kinase-related Cdc42-	1				
		binding kinase			(AF021936) myotonic		
		MRCK-beta (MRCK-			dystrophy kinase-related Cdc42-		
1776	A F00 10 3 4	beta) mRNA,	2- (2	2776152	binding kinase MRCK-beta	3 25	
1776	AF021936	complete cds	2e-62	2736153	[Rattus norvegicus] (AC004798) R31546_1 [Homo	3e-27	
1777	Y12059	H sapiens HUNKI mRNA	le-62	3184498		3e-09	
'''' 	112039	IUIVINA .	16-02	3104498	sapiens]	Je-UY	
		Human (clone E5.1)					
1 1		RNA-binding protein		•	sialidase - Actinomyces viscosus		
1778	L37368	mRNA, complete cds.	6e-63	477578	>gil141852	7.8	
	237300	mici vi t. complete cas.	<u> </u>		ZINC FINGER PROTEIN 83	,	
		Figure 2. Nucleotide			(ZINC FINGER PROTEIN		
		and translated protein			HPF1) >gi 106023 pir A32891		
1 1		sequences of HPF1, -			finger protein 1, placental -		
1779		2, and -9.	5e-63	1731443	human	3e-33	
		Homo sapiens					
		putative G protein-			(AL023288) hypothetical		
1780	AF095448	coupled receptor	2e-63	3116131	protein ·	4.6	
		II			•		
		Human transaldolase mRNA containing					
		transposable element.		l	(U63159) transaldolase [Mus	ĺ	
1781		complete cds	2e-63	1553119	musculus]	4e-18	
1/01	£19437	complete cus	26-03	1333113	PROSTASIN PRECURSOR	+C-19	
			1		precursor - human >gi 862305		
		Homo sapiens			(L41351) prostasin [Homo		
		prostasin mRNA,			[sapiens] >gi 1143194 (U33446)	Į	
1782		complete cds	Ie-63	2833277	prostasin [Homo sapiens]	6e-14	
1	Ĭ	Homo sapiens 10kD]				
		protein (BC10)			hypothetical protein K03H1.9 -		
1783	AF053470	mRNA, complete cds	6e-64	482237	Caenorhabditis elegans	0.029	

	Nearest	Neighbor (BlastN vs. G	ienbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ			1				
ĽD	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
						 	
				<u> </u>	(Z93390) Similar to FY VE zinc	 	
]				i	finger; cDNA EST yk265b4.5		
					comes from this gene; cDNA	1	
		!	•		EST yk359g9.5 comes from this	1	
				İ	gene; cDNA EST yk319c2.5		
	•				comes from this gene [Caenorhabditis elegans] zinc		
					finger; cDNA EST yk265b4.5		
					comes from this gene; cDNA		
					EST yk359g9.5 comes from this		
		Mouse mRNA for			gene; cDNA EST yk319c2.5		
		beta-1,4-			comes from this gene		
1784	D37791	galactosyltransferase	6e-64	3880102	[Caenorhabditis elegans]	3e-16	
		Mus musculus radical					
1705	A F016770	fringe (radical-fringe)		200.00	(U94350) radical fringe		
1785	AF015770	mRNA, complete cds H.sapiens flow-sorted	6e-64	2204355	precursor [Mus musculus]	le-36	
		chromosome 6			1		
		HindIII fragment.		·	1		
1786	Z79054	SC6pA21E11	2e-64	<none></none>	<none></none>	<none></none>	
		Homo supiens cytosolic selenium-					
ľ		dependent glutathione	Ī				
ŀ		peroxidase gene.			[
	i	complete cds, and	•		(U42580) A565R [Paramecium		
1787		rhoh 12 gene. 3' end.	le-64	2447063	bursaria Chlorella virus 1	8.8	
		H.sapiens LAG-3			(X14112) tegument protein		
1788	Y10211	gene, promoter region	7e-65	1944540	[human herpesvirus 1] .	2.3	
		Human lucaruma			·		
1789		Human lysozyme mRNA, complete cds.	2e-65	<none></none>	<none></none>	<none></none>	
		max. v.a., complete eds.	20-03		LIPASE MODULATOR	/INOINE>	
j	1				PRECURSOR (LIPASE		
ł	j		.		HELPER PROTEIN) .	ļ	
ļ		Homo sapiens SS-			>gi 480045 pir S36249 lipB		
Ī	The state of the s	A/Ro autoantigen 52			protein - Pseudomonas glumae		
		kda component gene.			>gi 49207 (X70354) helper	1	
1790		complete cds	2e-65	585401	protein	4.2	
		Homo sapiens RNA	i.				
		binding protein DEF- 3 mRNA, complete		ļ	(AFOCOSITA DAIA HISSISS		
1791		ods complete	2e-65	Į.	(AF069517) RNA binding	10.25	
	. 11 009317	.43	46-03	3212101	protein DEF-3 [Homo sapiens]	le-25	

<u> </u>	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		1		ivenest iveign	HOO! (Blasee Vs. 140h-Reddingant Pr	Oteins)	
D	ACCESSION	DESCRIPTION	P VALUE	1.CC=CC10.:	DE CONTROL		
-	TACCESSION		PVALUE	ACCESSION	DESCRIPTION	P VALUE	
_	 	Homo sapiens jerky gene product	 				
İ		homolog mRNA.	ŀ		(AE004215): 1		
1792	AF004715	complete cds	2e-65	2314829	(AF004715) jerky gene product homolog [Homo sapiens]		
				2314829	hypoxanthine (guanine)	2e-45	
ł	Į.				phosphoribosyltransferase - long	1	
i	j	C. longicaudatus hprt			tailed hamster		
		mRNA for			phosphoribosyltransferase		
1793	X59652	hypoxanthine	3e-66	631625	[Cricetulus longicaudatus]	6e-54	
1	i						
		Mus musculus radical					
1794	U94350	fringe precursor	2. 42	000.000	(U94350) radical fringe		
11.54	094330	mRNA, complete cds	3e-67	2204355	precursor [Mus musculus]	2e-33	
Ì	ſ	Mus musculus					
1	1	putative	- 1				
1 .	!	lysophosphatidic acid			(AF015811) putative		
]]		acyltransferase	j		lysophosphatidic acid		
1795	AF015811	mRNA, complete cds	3e-68	2317725	acyltransferase [Mus musculus]	7e-51	
		Cow					
	\	phosphoinositide-					
1796	J03137	specific phospholipase C	3e-69	224000	phospholipase C 154 (Bos		
1,70	703137	phosphotipase C	36-09	226908	taurus]	3c-25	
		Rattus norvegicus					
		putative peroxisomal	i				
]]		2.4-dienoyl-CoA	ŀ				
]]		reductase (DCR-			(AF044574) putative		
		AKL) mRNA,	i		peroxisomal 2,4-dienoyl-CoA	Ì	
1797	AF044574	complete cds	1e-69	4105269	reductase (Rattus norvegicus)	2e-33	
		Mus musculus				ľ	
		putative	ĺ				
1	l l	lysophosphatidic acid	.]		(AF015811) putative		
		acvltransferase	1		lysophosphatidic acid	J	
1798		mRNA, complete cds	4e-70	2317725	acyltransferase (Mus musculus)	3e-19	
		M.musculus mRNA	1		and the state of t	36-19	
		for desmoyokin.	1		desmoyokin - mouse (fragment)	- 1	
1799	X65157	partial	5e-74	109781	>gi 50675	9e-37	
- 1			- 1				
1800		Mus musculus mR.NA	a. 7.	202.0.5	(Z97207) B-IND1 protein [Mus		
.000	251201	or B-IND1 protein	2e-74	2231019	musculus]	6e-21	
	l	Gallus gallus zinc					
1		inger protein (Fzf-1)	1		(U27196) zinc finger protein		
1801		nRNA. complete cds.	6e-75	984814	[Gallus gallus] gallus]	2e-44	
	L			20.017	(Carres Entres)		

	Nearest	Neighbor (BlastN vs. (Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC			1			T. C. C. C. C. C. C. C. C. C. C. C. C. C.	
D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					70 KD WD-REPEAT TUMOR-	I VALUE	
		Rattus norvegicus			SPECIFIC ANTIGEN		
	1	mRNA for 70 kDa			>gi 2505957 gnl PID e353992		
		tumor specific			(Y15054) 70 kD tumor-specific		
1802	Y15054	antigen, partial	3e-77	3123027	antigen [Rattus norvegicus]	4 c- -12	
		M.musculus mRNA			- Interest (Italian)	40-42	
1	1	for desmoyokin,	1 .	•	desmoyokin - mouse (fragment)		
1803	X65157	partial	3e-79	109781	>gi 50675	9e-33	
						30 33	
1	l	Rattus norvegicus	İ .		cytokine inducible nuclear		
	İ	cardiac adriamycin	1		protein C193 - human		
1	I	responsive protein	1		>gi 793841 (X83703) nuclear		
1804	U50736	mRNA, complete cds	2e-84	1362781	protein [Homo sapiens]	7e-30	
i			•				
		Rattus norvegicus					
ı		thioredoxin reductase			·		
1 .		(TrxR2) mRNA,					
1	l	nuclear gene					
1	i	encoding					
1		mitochondrial			(AF072865) thioredoxin		
1805	AF072865	protein, complete cds	2e-84	3757888	reductase [Rattus norvegicus]	6e-43	
	٠	Rattus norvegicus	}		}		
		putative peroxisomal	j			ľ	
		2.4-dienoyl-CoA	í			ľ	
	j	reductase (DCR-	ı		(AF044574) putative	l	
1806	A FO 1 457 4	AKL) mRNA.	1		peroxisomal 2,4-dienoyl-CoA	j	
1300		complete cds	6e-85	4105269	reductase (Rattus norvegicus)	1e-41	
1 1		Rattus norvegicus	ł				
1807		Rabin3 mRNA,		44.000	(U19181) Rabin3 [Rattus		
1007	U19181	complete cds.	2e-87	624225	norvegicus]	2e-∔1	
		Mus musculus ninein	i		(7110212)		
1808		mRNA, complete cds.	le-91	1112066	(U40342) ninein [Mus		
		R.norvegicus mRNA	16-31	1113865	musculus] TROPOMYOSIN BETA 3,	2e-36	
! i		for cytosolic			1	1	
		resiniferatoxin-	ł		FIBROBLAST chicken	1	
1809		binding protein	4e-92	136077	>gi 515694 (M23082)	1	
		ding protein	70-72	130077	tropomyosin (Gallus gallus)	0.56	
l		Rattus norvegicus	ł			l l	
		outative peroxisomal	1				
		2.4-dienoyl-CoA	1			į	
l		reductase (DCR-	J		(AF044574) putative	- 1	
		AKL) mRNA.	1		•		
1810		complete cds	5e-93	4105269	peroxisomal 2.4-dienoyl-CoA	, , ,	
		vlus musculus EHF	30-23	+100-09	reductase [Rattus norvegicus]	1e-50	
- 1		Ehf) mRNA.	- 1		(A E035527) EWE (M		
1811		omplete cds	2e-95	3138930	(AF035527) EHF (Mus	2	
		ompiere cas		טנעטנונ	musculus]	2e-47	

	Nearest	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ			1		1001 (2) and 1 st. 14011-Redundant P	Toteris)		
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION			
-	1	1 3000	I VALUE	ACCESSION	DESCRIPTION	P VALUE		
	 	Cricetulus griseus	 			ļ <u> </u>		
		mRNA for			1			
ļ	1	Phosphatidylglycerop			(AB016930)			
		hosphate synthase,		•	Phosphatidylglycerophosphate	1		
1812	AB016930	complete cds	6e-96	4159682	synthase [Cricerulus griseus]	7- 41		
				1157002	Symmote (Cricetalus griseus)	7e-41		
	ł	Rattus norvegicus						
	ł	mRNA for atypical			(AB005549) atypical PKC	[
1	1	PKC specific binding			specific binding protein [Rattus			
1813	AB005549	protein, complete cds	7e -97	3868778	norvegicus]	3e-41		
1					polybromo I protein - chicken			
1		ŀ			chicken >gi 951231 (X90849)	f .		
1814	35000.40				polybromo I protein (Gallus			
1014	X90849	G.gailus PB i gene h-lamp-2=lysosome-	2e-97	2134381	gallus]	le-34		
		associated membrane	i		1			
1		protein-2 protein-2b	ļ					
		(LAMP2) mRNA,			1			
		alternatively spliced	l					
!		form h-lamp-2b,	Į.					
1815	S79873	complete cds.	3e-98	<none></none>	<none></none>	<none></none>		
		Mus musculus ACF7			T. C.	CHOINES		
		neural isoform 1						
		(mACF7) mRNA,			(U67204) ACF7 neural isoform]		
1816	U67203	partial cds	2e-98	1675224	2 [Mus musculus]	9e-39		
			i					
1 1	,	Rattus norvegicus	į		ELONGATION FACTOR G.			
·		nuclear-encoded	1		MITOCHONDRIAL	1		
		mitochondrial	1		PRECURSOR (MEF-G)	i		
1 1	l	elongation factor G	- 1		>gi 543383 pir S40780			
1817		mRNA, complete cds.	e-100	585084	translation elongation factor G.	2. 22		
		in the complete cus.		303004	mitochondrial - rat >gi 310102 spermatid perinuclear RNA-	2e-30		
				•	binding protein Spnr - mouse			
	ļ	M.musculus Spnr	İ		>gi 673454 (X84692) spermatid	j		
		mRNA for RNA	}		perinucleur RNA binding			
1818	X84692	binding protein	e-133	1363238	protein [Mus musculus]	5e-35		
		Rattus norvegicus			cytokine inducible nuclear	j		
1	I.	cardiae adriamycin			protein C193 - human	1		
1810		responsive protein			>gi 793841 (X83703) nuclear			
1819		nRNA, complete cds	e-113	1362781	protein [Homo sapiens]	2e-36		
1		HoxB9=Hox-2.5 mice. embryos,						
- 1		nRNA Partial, 786			Volume november 1			
1820		nkiya Partial. 700	0.107		HOMEOBOX PROTEIN HOX-	1		
	200033	···	e-107	1708355	B9 (HOX-2.5)	Se-37		

	Near	est Neighbor (BlastN vs	. Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant F	
SI	EQ				griedi (Siasur 13: Noil-Redundant F	roteins)
T.	D ACCESSI		P VALUE	ACCESSION	DESCRIPTION	P VALUE
		HoxB9=Hox-2.5				IF VALUE
		[mice, embryos, mRNA Partial, 786			WOMEO DOWN	
18:	21 S66855	nt]	e-108	1708355	HOMEOBOX PROTEIN HOX B9 (HOX-2.5)	4e-37
		Rattus norvegicus n tomosyn mRNA,	n-			46-37
18:	22 U92072	complete cds	e-102	3790389	(U92072) m-tomosyn [Rattus norvegicus]	
- 1	1			0.0000	KINESIN-LIKE PROTEIN	2e-38
	1		1		KIF1B mouse	
- 1		Mouse mRNA for	ı		>gi 407339 gn1 PID d1005029	
1,00		kinesin-like protein	1	i	(D17577) Kiflb (Mus]
182	23 D17577	(Kif1b), complete co	is e-129	2497524	musculus)	2e-39
		Mus musculus SDPS			(AE0(248))	
182	4 AF062484	mRNA, complete cd	s e -122	3126981.	(AF062484) SDP8 [Mus musculus]	£ . 10
	1				THE POPULATION (ATOMS)	5e-40
			ĺ		(H3.3Q) histone H3.3 - fruit fly	
1	1			İ	(Drosophila melanogaster)	i
	İ				histone H3.3B - chicken	i
1	1		1 1		>gi 2119023 pir S61218 histone	ł
1		j	1		H3.3 - fruit fly (Drosophila	1
1					hydei) I-136) [Oryctolagus	1
1	Į.		1 1		cuniculus] >gi 8046 (X53822)	- 1
1	1		! !		Histone H3.3Q gene product	Į.
1	1		1 1		[Drosophila melanogaster]	1
1	1		1 . 1		>gi[5[198 gallus] >gi[16] 190	ł
1			1 1		(M17876) histone H3 [Spisula	
1	ł		1 1		solidissima] >gi 211853	i i
1	l	i	1 1		(M11393) histone 3.3 [Gallus	1
l I	l		1		gallus] >gi 306848 (M11354)	1
]	1	1			H3.3 histone [Homo sapiens]	ľ
1		ľ	1		melanogaster] >gi 963031	
l i		R.norvegicus mRNA			(X81205) histone H3.3 H3.3A	1
1825	X73683	for histone H3.3	e-109	122075	variant [Drosophila	
		Mus musculus ACF7		1220/3	melanogaster) musculus)	2e-40
		neural isoform 1	1			ĺ
	_	(mACF7) mRNA,			(U67204) ACF7 neural isoform	
1826	U67203	partial cds	e-102	1675224	2 [Mus musculus]	20 10
· [KINESIN-LIKE PROTEIN	2e-40
ł			1		KIF1B mouse	
		Mouse mRNA for	- 1		>gi 407339¦gnl PID d1005029	1
1827		kinesin-like protein	1		(D17577) K:flb [Mus	
1027	D17577	(Kiflb), complete cds	e-131		musculus]	7e-42

1	Neares	st Neighbor (BlastN vs. C	ienhank)	None N	The CDL of	
SE			T T	Nearest Neig	hbor (BlastX vs. Non-Redundant F	roteins)
П	7 [N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-						1
- 1	1	Cricetulus griseus	ļ			
		mRNA for		j		i i
	ď	Phosphatidylglycerop		i	(AB016930)	1
182	8 AB016930	hosphate synthase.		1	Phosphatidylglycerophosphate	1
1.01	O AD010930	complete cds	e-131	4159682	synthase [Cricetulus griseus]	3e-43
	1	Mus musculus SKD3				
182	9 U09874	mRNA, complete cds.	e-122	7402776	SKD3 PROTEIN SKD3 (Mus	1 1
		mad vi i, complete eds.	C-122	2493735	musculus]	7e-48
- 1	i	C.familiaris mRNA			(X99145) overexpressed in thyroid tissue after TSH	1
183	0 X99145	for C3VS protein	.e-110	1429314	stimulation [Canis familiaris]	
		P.walti mRNA for		. 127514		2e-49
		rmp associated protein	٠.		(X99836) rap55 [Pleurodeles	
183	1 X99836	55	e-106	4200286	walti]	2e-50
						26-30
1		Mus musculus SH3				
	İ	domain-containing			(AF077003) SH3 domain-	! [
1832	AF077003	adapter protein			containing adapter protein;	
1832	AF077003	mRNA, complete cds	e-121	3550240	CD2AP	3e-51
	ł	Mus musculus strain	,			
	1	C57BL/6 zinc finger	J			
1	i	protein 106 (Zfp106)]	
1		mRNA, H3a-a allele.	- 1		(AFOCOOLG)	1
1833	AF060246	complete cds	c-118	3372657	(AF060246) zinc finger protein	·
				3372037	106 [Mus musculus] TRANSLOCON-	1e-52
1	1		1		ASSOCIATED PROTEIN.	i
1]				GAMMA SUBUNIT (TRAP-	
	f		1		GAMMA) (SIGNAL	ł
1			ľ		SEQUENCE RECEPTOR	
1	[1		GAMMA SUBUNIT) (SSR-	- 1
1		D	- 1		GAMMA)	j
		R.norvegicus mRNA	· 1		>gi 423185 pir S33294	
1834		for TRAP-complex			translocon-associated protein	1
	214030	gamma subunit.	e-120	1174453	gamma chain - rat norvegicus]	7e-5∔
		Mus musculus SH3	j		1	
	ı	domain-containing	. 1		(AE077002) SU2 4	Ī
		adapter protein	1		(AF077003) SH3 domain-	ı
1835		mRNA, complete cds	e-132		containing adapter protein; CD2AP	
	T				(L20427)	5e-54
[Rattus norvegicus	1		dihydroxypolyprenylbenzoate	
		dihydroxypolyprenylb	ŀ		methyltransferase	
		enzoate	1		dihydroxypolyprenylbenzoate	j
1024		nethyltransferase		ļ	methyltransferase [Rattus	Ì
1836	L20427 r	nRNA, complete cds	e-116		norvegicus	4e-56

	Near	st Neighbor (BlastN vs	. Genbank)	Nearest Vei	ghbor (BlastX vs. Non-Redundant	0
SE	Q			- Carest : te.	gilosi (Biasot Vs. 14011-Redundant	Proteins)
	ACCESSIO	ON DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
<u> </u>					PROTEIN TSG24 (MEIOTIC	IP VALUE
- 1	1				CHECK POINT	
		M.musculus mRNA		i	REGULATOR)	
18:	37 X80169	for 200 kD protein	e-122	1717793	>gi 1083553 pir A55117 tsg24	1
					P 5:11005555 PH A 55117 CS 24	2e-56
- 1	i	Rattus norvegicus				1
-		CTP:phosphoethano	ol [1		ļ
ŀ	ł	amine		i	(AF080568)	
		Cytidylyltransferase	[1	CTP:phosphoethanolamine	1
183	8 AF080568	mRNA, complete co	ls e-119	3396102	cytidylyltransferase	6.50
l	ı				(X99145) overexpressed in	6e-58
1.00		C familiaris mRNA			thyroid tissue after TSH	
183	9 X99145	for C3VS protein	e-121	1429314	stimulation [Canis familiaris]	2e-58
-					(Carro ratification)	26-38
ļ	ł	Pan troglodytes brea	st	f	·	1 1
1	1	and ovarian cancer				1 1
1	1	susceptibility			(AF005068) breast and ovarian	
1,04		(BRCA1) gene,	1		cancer susceptibility protein	1 1
1840	AF019075	17	e-145	2218154	splice variant [Homo sapiens]	1e-58
1841	1155040	Bos taurus myosin X			(U55042) myosin X [Bos	10-30
1041	U55042	complete cds	e-122	1755049	taurus]	le-61
	1	Mus musculus mRN	.] [13.01
1	j	for poly(ADP-ribose)				1 1
1842	AJ007780	polymerase-2	1 1		(AF072521) poly-(ADPribosyl)-	1 1
	1.007700	potymerase-2	e-119	3283975	transferase homolog PARP	4e-62
1	ļ	Rattus norvegicus	1 1			
		thioredoxin reductase	1 1		1	1
1	j	(TrxR2) mRNA.	1 1		1	
	•	nuclear gene	1 1		1	1
İ		encoding	1 1		1	i
1		mitochondrial	1			İ
1843	AF072865	protein, complete cds	e-105	3757888	(AF072865) thioredoxin	
		Bos taurus myosin X.	57105	3737000	reductase [Rattus norvegicus]	3e-62
1844	U55042	complete cds	e-121	1755049	(U55042) myosin X [Bos	
		Mouse E46 mRNA	- 121	1133049	taurus]	le-62
1845	X61506	for E46 protein	e-139	114909	BRAIN BROTTING	
- 1				114707	BRAIN PROTEIN E46 GUANINE NUCLEOTIDE	9e-67
ľ			1		BINDING PROTEIN, ALPHA-	1
- 1	ſ		1		14 SUBUNIT (GL1)	1
ŀ	ł		i		>gi 108711 pir A40891 GTP-	1
]		Bovine mRNA for			binding protein GL1 alpha chain	1
		GTP-binding protein	1		- bovine protein, alpha-subunit	ľ
1846		alpha-subunit	e-148	585174 .	[Bos taurus]	20.60
		Mus musculus			(2e-69
10.7		B6CBA Lisch7			(AC002128) Lisch7 [Homo	1
1847	U49507	mRNA, partial cds.	e-140		sapiens]	2e-7∔
						-6-/4

Table 4

	Nearest N	leighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant F	Democratic V
SEC				 	Show i Blaset vs. : toll-Reddildalit i	Toteins)
ΙD	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
 	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
2	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
3	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
4	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
5	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
6	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
7	<none></none>	<none></none>	<none></none>	<none></none>	· <none></none>	<none></none>
8	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
9	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
10	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
[1]	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
12	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
13	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
[4	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
15	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
16	<none></none>	<none></none>	<none></none>	<none></none>		<none></none>
17	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
18	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
19	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
20	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
21	<none></none>	<none></none>	<none></none>	<none></none>		<none></none>
22	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
			1.0.12	CHOINES	<none> tMDC I protein - crab-eating</none>	<none></none>
. 23	<none></none>	<none></none>	<none></none>	1079469		
			a.io.i.es	10/2409	(AB011138) KIAA0566 protein	9.3
24	<none></none>	<none></none>	<none></none>	3043656	[Homo sapiens]	
				30 13030	[[Homo sapiens]	9.3
- 1]		notossium ab annul manui a B 155	1
25	<none></none>	<none></none>	<none></none>	112175	potassium channel protein RK5	2.6
			W.O.L.	1121/3	rat protein [Rattus norvegicus]	8.6
26	<none></none>	<none></none>	<none></none>	3769624	(AF091565) olfactory receptor	1
		41.01.2	KNONE	3709024	[Rattus norvegicus]	7.2
27	<none></none>	<none></none>	<none></none>	3876443	(Z81517) F28B1.6	
			VIIIONE -	3070443	[Caenorhabditis elegans]	7. l
28	<none></none>	<none></none>	<none></none>	2221164	(AB001684) ORF249 [Chlorella	
		2.01.0	CITOINES	2224464	vulgaris]	6.9
- 1			J		(U67940) ORFveg106; random	
29	<none></none>	<none></none>	<none></none>	1610707	cDNA sequence [Dictyostelium]	
		V. TOTTE	CIVOIVE	1519707	discoideum]	6.7
30	<none></none>	<none></none>	<none></none>	227.101	protein kinase C II [Xenopus	
		- CIVOINE	CHOILES	227491	laevis]	6.7
31	<none></none>	<none></none>	NONE	(20.77	C50C3.4 protein -	ł
- 	41.01.12	CHOILES	<none></none>	630575	Caenorhabditis elegans	6.0
- 1	·	J	j		35 KD PROTEIN IN RNA2	ł
- 1			Ì		clover necrotic mosaic virus	1
- 1		ĺ	j		>gi 61466 (X08021) ORF for 35	ı
1	1 *	l	ſ		kDa polypeptide (AA 1-317)	i
32	NONE-	2,0712			[Red clover necrotic mosaic	1
	<none></none>	<none></none>	<none></none>	137290	virus]	6.0

-	Neare	st Neighbor (BlastN v.	s. Genbank)	Nearest N.	eighbor (DL - V	
SE	Q			TACATEST 146	eighbor (BlastX vs. Non-Redundant F	roteins)
IL	ACCESSIO	DESCRIPTION	PVALU	ACCESSION	N DESCRIPTION	P VALU
33	<none></none>	<none></none>	NONE	-	(X16711) pid:g30041 [Homo	1
		- CANOLINE	<none></none>	30041	sapiens]	5.9
-34	<none></none>	<none></none>	<none></none>	2402505	CELL DIVISION PROTEIN	
35	<none></none>	<none></none>	<none></none>	2175505	FTSW	5.7
			- VIONE	1001430	(D63999) hypothetical protein	5.7
36	<none></none>	<none></none>	<none></none>	3182918	NITROGEN REGULATORY	
	1			3102718	PROTEIN AREA MITOCHONDRIAL	5.2
			1	1	RIBOSOMAL PROTEIN SS	
	1	1	İ	1	Emericella nidulans	ĺ
	ı		Ī	1.	mitochondrion (SGC3)	
37	ANONT			•	>gi 12709 nidulans] >gi 472822	·
	<none></none>	<none></none>	<none></none>	140011	(J01390) unknown protein	4.3
	I				(AL034393) predicted using	4.3
	i	1	j		Genefinder; similar to WD	
			}	1	domain, G-beta repeat; cDNA	
			1	1	EST yk362f7.5 comes from this	
			1		gene; cDNA EST yk362f7.3	
38	<none></none>	<none></none>			comes from this gene	
	THORNE	CITOINES	<none></none>	3979943	[Caenorhabditis elegans]	4.0
39	<none></none>	<none></none>	<none></none>		(U31329) polyketide synthase	
		Q.O.L.	KNOWES	950203	[Aspergillus terreus]	3.3
1			1 1			
	1		1 1		(AL031530) hypothetical zinc	
40	<none></none>	<none></none>	<none></none>	3560232	finger protein	
- 1				3300232	[Schizosaccharomyces pombe]	3.0
1			1		AXONEME-ASSOCIATED	
41	<none></none>	<none></none>	<none></none>	730071	PROTEIN MST101(1) product	f
	1			7,5007,1	[Drosophila hydei] HYPOTHETICAL 21.7 KD	2.6
- 1	ł				PROTEIN IN INTE-PIN	1
- 1	. 1	ŀ			INTERGENIC REGION	ł
- 1	.	'	- 1		>gi 1787402 (AE000214) orf,]
.2	<none></none>	2102			hypothetical protein	
-	-MONES	<none></none>	<none></none>	2506641	[Escherichia coli]	2.5
3	<none></none>	-NONT:			(AF071556) anthranilate	2.5
+		<none></none>	<none></none>	3511232	dioxygenase large subunit	2.4.
	1		1			
ı	- 1		- 1		(U43139) envelope glycoprotein	ł
1	<none></none>	<none></none>	<none></none>		gp120 (Human	
			ZITOMES	1150900	immunodeficiency virus type 1]	1.9
	1	1	J		(Z75536) similar to dynein	
-	I	İ	ł		heavy chain; cDNA EST	1
1	1		ſ		EMBL:D27549 comes from this	
1	İ		ľ		gene; cDNA EST]
	<none></none>	<none></none>	<none></none>	3876099	EMBL:D34859 comes from this	
			24		gene [Caenorhabditis elegans]	1.4

	17	earest (Veighbor (BlastN	vs. Genbank)	Nearest N	leighbor (BlastX vs. Non-Redundar	
	EQ ID ACCES	SION	DESCRIPTIO	N P VALU			
	46 <noi< td=""><td></td><td></td><td></td><td></td><td>(AL032647) predicted using</td><td>P VALU</td></noi<>					(AL032647) predicted using	P VALU
	46 <no< td=""><td>VE></td><td><none></none></td><td><none< td=""><td>> 3881150</td><td>Genetinder</td><td></td></none<></td></no<>	VE>	<none></none>	<none< td=""><td>> 3881150</td><td>Genetinder</td><td></td></none<>	> 3881150	Genetinder	
1	1	- 1		1		COLANIC ACID CAPSULA	1.4
	- 1	1		1	ł	BIOSYNTHESIS	LK
1	j	- 1		1	1	ACTIVATION PROTEIN A	1
4	7 <non< td=""><td>re.</td><td>-NONE</td><td>- 1</td><td>1</td><td>>gi 95605 pir S 17701 rcsA</td><td>1</td></non<>	re.	-NONE	- 1	1	>gi 95605 pir S 17701 rcsA	1
		-	<none></none>	<none></none>	132200	protein	1 , ,
4	8 <non< td=""><td>E> </td><td> <none></none></td><td></td><td></td><td>(U61380) germination protein</td><td>1.1</td></non<>	E>	<none></none>			(U61380) germination protein	1.1
			THORES	<none></none>	2204286	[[Bacillus megaterium]]	1.0
	1			i		HTPOTHETICAL TLA KD	1.0
	1				1	PROTEIN IN FOXI-KEXI	- { - ·
	1				1	INTERGENIC REGION	
	1	1			1	>gi 2132566 pir S64222	-1
	1	- 1		- 1		probable membrane protein	1
	· I				j	YGL204c - yeast	1
	ı	- 1		ı	1	(Saccharomyces cerevisiae)	1
	1			1	1	>gi 1322838 gnl PID e243803	1
49	<none< td=""><td>></td><td><none></none></td><td><none></none></td><td>177205 -</td><td>(Z72726) ORF YGL204c</td><td>ſ</td></none<>	>	<none></none>	<none></none>	177205 -	(Z72726) ORF YGL204c	ſ
	1	T		11.0.112	1723955	[Saccharomyces cerevisiae]	0.84
	1			1	1	(AJ006514) prolipoprotein	
50	<none< td=""><td><u> </u></td><td><none></none></td><td><none></none></td><td>3201564</td><td>diacylglyceryl transferase</td><td>1 1</td></none<>	<u> </u>	<none></none>	<none></none>	3201564	diacylglyceryl transferase	1 1
51		1			3201304	[Vibrio cholerae]	0.31
<u> </u>	<none< td=""><td>`</td><td><none></none></td><td><none></none></td><td>2808721</td><td>(AL021428) hypothetical protein Rv0064</td><td></td></none<>	`	<none></none>	<none></none>	2808721	(AL021428) hypothetical protein Rv0064	
52	CNIONIE.					(U17986) GABA/noradrenaline	0.27
	<none:< td=""><td>-</td><td><none></none></td><td><none></none></td><td>602434</td><td>transporter [Homo sapiens]</td><td></td></none:<>	-	<none></none>	<none></none>	602434	transporter [Homo sapiens]	
	1		·			(AF076184) cytosolic sorting	0.13
53	<none></none>		-MONTE	1 1		protein PACS-1b [Rattus	1 1
_		-	<none></none>	<none></none>	3347955	norvegicus]	0.12
						(U53344) coded for by C.	0.12
				1		elegans cDNA yk92b4.5: coded	1 1
				, ,		for by C. elegans cDNA	1
- 1		1		1 1		yk73a1.5; coded for by C.	
- [l			1	•	elegans cDNA.yk102e9.5;	
- 1		i		1		coded for by C. elegans cDNA	į
- 1		1	1	J		yk71c8.5; coded for by C.	l l
. 1		1				elegans cDNA yk66d11.5;	ł
-	<none></none>		<none></none>	<none></none>	1255887	coded for by C. elegans cDNA	i
					1433887	yk66c3	0.074
- 1		1 .	1	1		Bkm-like sex-determining	
			1	- 1		region hypothetical protein	. 1
-1	<none></none>	ļ	<none></none>	<none></none>	103076	CS314 - fruit fly (Drosophila melanogaster)	f
\neg					103070	IMCDDOugstee)	0.00-
T	<none></none>	l	<none></none>			Ras inhibitor (clone JC265) -	0.003

8	Nearest	Neighbor (BlastN vs. (Genbank)	Nearest Neigh	nbor (BlastX vs. Non-Redundant Pi	roteins)
SEQ						
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	PVALUE
		† 	1		Bkm-like sex-determining	1
			 		region hypothetical protein	
			1		CS314 - fruit fly (Drosophila	
57	<none></none>	<none></none>	<none></none>	103076	melanogaster)	2c-04
					(AF038604) contains similarity	
1			ŀ	<u> </u>	to Drosophila ovarian tumor	
i l		1			locus protein (GB:X13693)	
58	<none></none>	<none></none>	' <none></none>	2702370	[Caenorhabditis elegans]	6e-05
					(AL033501) phox domain	
59	<none></none>	<none></none>	<none></none>	3859713	protein [Candida albicans]	3e-05
		· · · · · · · · · · · · · · · · · · ·			(AF003386) F59E12.5 gene	
		'			product [Caenorhabditis	
60	<none></none>	<none></none>	<none></none>	2088839	elegans]	2e-08
7					GC-RICH SEQUENCE DNA-	
		`	1	ł	BINDING FACTOR GCF -	
			}		human >gi 179412 (M29204)	
i 1					DNA-binding factor (Homo	
61	<none></none>	<none></none>	<none></none>	121059	sapiens]	4e-09
•			Į i		(Z81490) similar to WD	
]		domain, G-beta repeats (2	
Ì			Ī		domains); cDNA EST	
]		EMBL:T00482 comes from this	
					gene; cDNA EST	
					EMBL:T00923 comes from this	1 1
					gene; cDNA EST yk449d4.3	
					comes from this gene; cDNA	
62	<none></none>	ANONIES	NONT	2075246	EST yk449d4.5 comes from this	
02	CIVOIVES	<none></none>	<none></none>	3875246	gen	9e-24
				\ ////	(U64857) No definition line	
63	<none></none>	<none></none>	<none></none>	1465834	found [Caenorhabditis elegans]	9e-28
	VIVOIVE -	CHOILE	CHOILES	1403034	(AB014561) KIAA0661 protein	9C-26
64	<none></none>	<none></none>	<none></none>	3327136	[Homo sapiens]	1e-29
			3		(Z66521) similar to	 ``
- 1	ľ		}]		mitochondrial RNA splicing	
	ļ		ļ i		MSR4 like protein; cDNA EST	
1					EMBL:C09217 comes from this	
65	<none></none>	<none></none>	<none></none>	3880433	gene [Caenorhabditis elegans]	8e-31
		Rat annexin V gene,				
66		exon7 and exon8	5.0	<none></none>	<none></none>	<none></none>
					(U41278) coded for by C.	
- 1	1	•			elegans cDNA yk79g8.5; coded	
j	l				for by C. elegans cDNA	ı
- 1	j		[cm10c8; coded for by C. elegans	
- 1		Homo sapiens	l		cDNA yk79g8.3; similar to]
- 1		(subclone H8 2_d11	1		leucine-rich repeats found in	
		from P1 35 H5 C8)	_		many proteins (Caenorhabditis	
67	L35679	DNA sequence.	5.0	1086902	elegans]	6.6

	Nearest	Neighbor (BlastN vs. (Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ			1		VS. Non-Redundant Pr	oteins)	
ID	ACCESSION	DESCRIPTION	P VALUE	A COPOCYON			
-	T. CCZCBIOI	IHIV-1 strain BX220	T VALUE	ACCESSION	DESCRIPTION	P VALUE	
├		from USA, envelope	 				
ŀ		glycoprotein C2V3	j				
	[region (env) gene,	[]			
68	U90184	partial cds	50	1007070	(Z71986) convicilin precursor		
	0,0104	Human myosin VIIa	5.0	1297070	[Vicia narbonensis]	6.6	
		(MYO7A) gene, 5'			(AE000535) L-lactate permease	l	
69	U61465	exon 37	5.0	2313225	(lctP) [Helicobacter pylori 26695]	5.0	
			5.0	2313223	(AF064869) brain-enriched	5.0	
		Homo sapiens			guanylate kinase-associated		
		periplakin (PPL)		i	protein 2; BEGA2 [Rattus	ļ	
70	AF013717	mRNA, partial cds	5.0	3719238	norvegicus)	3.8	
		Soybean mRNA for			nor regicus;	3.6	
71	X58245	HMG-1 like protein	5.0	2995363	(AL022245) biotin synthase	0.99	
		Frasera paniculata					
		tRNA-Leu (tmL)					
		gene, intron.			(AC004411) putative		
72	AF102425	chloroplast sequence	4.9	3522050	pectinesterase (Arabidopsis		
	12 102 423	emoropiasi sequence	4.9	3522958	(Z81494) CDNA EST	6.4	
					EMBL:D27474 comes from this		
					gene; cDNA EST		
]		EMBL:D27473 comes from this		
- 1					gene; cDNA EST		
					EMBL:T00471 comes from this		
					gene; cDNA EST		
· .					EMBL:D34192 comes from this		
<i>^</i>		H.sapiens	ŀ		gene; cDNA EST	j	
		PTP1C/HCP-variant	.i		EMBL:D37241 comes from this	į	
73	X82817	gene	4.9	3875514	gene;	2.8	
- [-	•	1	1	
	j	. 1	Ţ		(AL031765) I-	1	
I	-		· I	• 💉	evidence=predicted by content;		
			ſ		1-method=genefinder;084; 1-		
	1	1	1		method_score=31.96; 1-	l l	
					evidence_end; 2-	1	
					evidence=predicted by match; 2-	Į.	
- 1	1	<u> </u>			match_accession=SPTREMBL:	ľ	
J		Mus musculus brain	į.		Q93319; 2-		
		atty acid-binding			match_description=HYPOTHE		
74	U04827 F	rotein	4.9	3676132	TICAL PROTEIN C33A11.2.;	2e-09	
ĺ		Veospora hughesi	1	•			
- 1		train NE1 internal				İ	
		ranscribed spacer 1,	ļ				
75		omplete sequence	4.8	<none></none>	<none></none>	AYON'E.	
				31101127	<none></none>	<none></none>	

		Nen	est Neighbor (BlastN vs				
	SE		est reignbor (Blastin vs	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundan	t Proteins)
		-		İ			
	H	ACCESSI			ACCESSION	DESCRIPTION	P VALUE
	70	3,0000	M.musculus MFH-	I			I. VALUE
		5 Y08222		4.8	<none></none>	<none></none>	<none></none>
		Í	Borrelia burgdorfer	i		1	KINOINES
	1	1	left chromosomal	1		·	1
	1 77	AJ22447	subtelomeric region	1		(AJ236702) HMR1 protein	i
	 ''	AJ2244/	(pfpB gene) Mus musculus LAF	4.8	4218141	[Antirrhinum majus]	8.3
	ı		putative membrane	1			- "" -
	1	1		1	1	,	
	l .	į.	protein (KRAG)			(AP000006) 367aa long	
	78	U02486	gene, exon 3 and	1		hypothetical protein	
	 	002480	Rat mRNA for	4.8	3258103	[Pyrococcus horikoshii]	2.7
		1	peptide/histidine				
		1	transporter, complete		ł	İ]]
	79	AB00028	0 cds		İ	(M29067) unknown protein	
		1 200020	o cus	4.8	806317	[Saccharomyces cerevisiae]	0.001
			A.cepa mitochondria	1			
		1	gene for NADH	"		i .	1
-		i	dehydrogenase	1	ł		} }
1		1	subunit 3 and		i	Į.	1 1
1		i .	ribosomal protein	1	1	1	1 1
L	_80	Z49771	S12	4.5	MONE		
ſ			Mouse IgG receptor	7.5	<none></none>	<none></none>	<none></none>
1			(beta-Fc-gamma-RII)			1	1
ı		i	gene, exons 6 and 7,	1 1			1 1
1			clones lambda-	1 1		1	1 1
L	81	M63494	Fc(3.2,93).	4.3	<none></none>	<none></none>	
ı		İ			4.ONES	<none></none>	<none></none>
ı		İ		1 1		(AF099000) No definition line	1 1
F	82	Z14035	S.pombe carl gene	2.0	3790665	found [Caenorhabditis elegans]	1 , 2
			Rhodococcus			(AL021687) putative protein	1.2
l	i		erythropolis ThcA	i i		[Arabidopsis thaliana]	1 1
l	· j		(thcA) gene, complete			>gi 2832633 gnl PID e1249651	1.
l	02	•••	cds; and unknown			(AL021711) putative protein	i
H	83	U17129	genes	2.0	2828280	[Arabidopsis thaliana]	2e-26
	- 1		Plasmodium			, , , , , , , , , , , , , , , , , , , ,	26-20
			falciparum	1		i	
	j		chromosome 2,	1			
			section 23 of 73 of	1			1
	84	A E00 1207	the complete			(AL031177) dJ889M15.3 (novel	
_	"	AE001386	sequence	2.0	4176500	protein)	9e-59
,	85	1170202	Human clone 23734				
	 -	U79292	mRNA sequence	1.9	<none></none>	<none></none>	<none></none>
	- }		Chloroplast Euglena	1			
	- 1		gracilis gene coding	1		·	
5	36	V00159	for the 5S and 16S rRNA.				
`		. 00137	INITA.	1.9	<none></none>	<none></none>	<none></none>

	Neare	st Neighbor (BlastN vs.	Genbank)	Negrect No	ighbor (Plane)	
SE	0			ivearest ive	ighbor (BlastX vs. Non-Redundant	Proteins)
II			P VALUE	ACCESSION	N DESCRIPTION	DVALE
-		Xenopus laevis XL-		T		P VALU
		INCENP (XL-		 		
	. 1	INCENP) mRNA,	1	i		
87	U95094	complete cds	1.9	<none></none>		
		H.salinarium TATA		SHORES	<none></none>	<none></none>
ł	1	box-binding protein	· [i	1	
88	X93206	genes and ORFs	1.9	<none></none>	1	ł
ĺ			 	SHOWES	<none></none>	<none></none>
	1	Caenorhabditis	ſ	•	1	
	1	elegans programmed]	1	1	ł
	1 .	cell death specifier	1	1		1
	I	(ces-2) gene,	1	1 .		1 .
89	U60979	complete cds	1			
	1	C. tentans ORF's (A-	1.9	<none></none>	<none></none>	_ <none></none>
90	X56272	E) for hemoglobin	1			
		Homo sapiens DNA	1.9	<none></none>	<none></none>	<none></none>
	1	sequence, repeat				
91	L22383	region.				1
		Hirudo medicinalis	1.9	<none></none>	<none></none>	<none></none>
	Ì	neuron-specific				
	i	protein mRNA,				
92	U82814	complete cds			(AF094531) immunoglobulin	1 1
		Haplomitrium	1.9	3822533	heavy chain precursor	2.0
į		hookeri 18S rRNA	i			
	[gene, partial	1		i	
93	U18504	sequence.			hypothetical protein 6 - fowlpox	1
		Pseudomonas stutzeri	1.9	1083969	virus virus]	2.0
		nosDFY genes	1			
		involved in copper	i			1
94	X53676	processing				1
		processing	1.9	2980781	(AL022198) putative protein	0.70
1		Dictyostelium]			
J		discoideum multidrug	ł			
- 1		resistance	ŀ	•		1
- 1		transporter/Ser	ı		(Z49130) cDNA EST	f
- 1		protease (tagC)			yk486b9.3 comes from this	ł
5					gene; cDNA EST yk486b9.5	
_		mRNA, complete cds. Human putative G-	1.9	3879530	comes from this gene	6e-05
		protein-coupled	1			
- 1		receptor (GPR17)	I		(Z75550) similar to cell division	į į
6					control protein [Caenorhabditis	ł
十		gene, complete cds	1.9	3880034	elegans]	7e-1↓
		Sus scrofa lactoferrin	j			
- 1		nRNA, complete cds.		•		1
1		:: gb I2\$421 I2\$421	i	i		I.
-			J		}	1
,		sequence 5 from atent US 5571691	I			I
/			1.8	<none></none>		

	Near	est Neighbor (BlastN vs. (Genbank)	Nearest Ne	ighbor (BlastX vs. Non-Redunda	
SE	Q				Agricol (Blastit Vs. 14011-Redundar	rt Proteins)
1I	ACCESSION	ON DESCRIPTION	P VALUE	ACCESSION	N DESCRIPTION	
<u> </u>		S.iniae lctP & lctO			DESCRIPTION	P VALU
98	Y07622	genes and ORF1	1.8	<none></none>	NONE	
1				***************************************	<none></none>	<none< td=""></none<>
1	1				Í	
1	1	Mouse myristoylated		İ		ľ
ł		alanine-rich C-kinase				[
1 00	1	substrate (MARCKS)		[j
99	M60474	mRNA, complete cds.	1.8	<none></none>	<none></none>	ANIONIE
1	V. 222.	Homo sapiens FGFR-			T.O.	<none:< td=""></none:<>
100	Y13901	4 gene	1.8	<none></none>	<none></none>	-NONTE
1	I	Human Down			4.0.1	<none:< td=""></none:<>
	Į.	Syndrome region of		• .	1	1 .
101	7744400	chromosome 21,		,	1	ſ
101	U44400	clone A31D6-1D6.	1.8	<none></none>	<none></none>	<none></none>
l	ſ					CIVONES
ł	j	Ruminococcus albus	1		0	
	l	beta-glucosidase				
102	U92808	(gluA) mRNA,	ľ		1	
102	092808	complete cds Candida albicans	1.8	<none></none>	<none></none>	<none></none>
	1	argininosuccinate	1			1
	İ	lyase (ARG4) gene,	ł		·	Ì
103	L25051	complete cds.	1			
		Helicobacter pylori	1.8	<none></none>	<none></none>	<none></none>
		26695 section 24 of				
		134 of the complete	1		1	
104	AE000546	genome	1.8	21017		
		Xenopus laevis major	1.8	<none></none>	<none></none>	<none></none>
- 1		beta-globin gene,	ı			
105	J00978	complete cds.	1.8	-NONE.	4	
		Human		<none></none>	<none></none>	<none></none>
		immunodeficiency	1			
- 1		virus type 1 isolate			1	1 1
		JW95-5, vpr gene,	1.			1 . 1
106	U41716	complete cds.	1.8	<none></none>	<none></none>	
		G.gallus mRNA for			CNONES	<none></none>
107	X66286	tensin	1.8	<none></none>	<none></none>	NONE
- 1		Xenopus calbindin			- STONES	<none></none>
		D28k mRNA,	· · · · · · · · · · · · · · · · · · ·		ļ	1 1
108		complete cds	1.8	<none></none>	<none></none>	<none></none>
.09		rabbit embryonic beta-	I			ZVOIAE2
· • ·	J00664	4-globin gene.	1.8	<none></none>	<none></none>	<none></none>
1	j,	Uuman see	1			11.01.12
Į		Human erg protein (cts-related gene)	l			1 1
10		nRNA, complete cds.	1		(AE000693) hypothetical	1
		mara. complete cas.	1.8	2983160	protein [Aquifex acolicus]	7.7

_	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Nois	phor (Rigety up No - P. I	
SE	Q				hbor (BlastX vs. Non-Redundant Pr	oteins)
ID	ACCESSIO		P VALUE	ACCESSION	DESCRIPTION	P VALU
		Rat troponin T			(Z46595) incomplete interleukin	1
111	M80829	cardiac isoform gene	1		11 receptor isoform [Homo	
	1/180829	complete cds	1.8	999450	sapiens]	7.3
112	D27867	Cyprinus carpio c- myc gene for c-Myc,		·	BRANCHED-CHAIN AMINO ACID TRANSPORT SYSTEM CARRIER PROTEIN (BRANCHED-CHAIN AMINO ACID UPTAKE CARRIER) >gi 1075007 pir D64056 membrane-associated component, branched amino acid transport system (brnQ) homolog - Haemophilus influenzae (strain Rd KW20) system II carrier protein (brnQ)	
112	D37887	complete cds Homo sapiens G	1.8	3023408	[Haemophilus influenzae Rd]	7.2
13	AF019765	protein-coupled receptor kinase 1 and G protein-coupled receptor kinase 1b (GRK1) gene, alternatively spliced, axternative exon 6, exon 7, and partial cds	1.8	498643	(U10270) G-box binding factor 1 [Zea mays]	7.2
4	AF025967	Helicobacter pylori J166 virulence regulon transcriptional activator homolog gene, partial eds, strain-specific genomic sequence B2	1.8	3850108	(AL033388) putative calcium- transporting atpase [Schizosaccharomyces pombe] PROBABLE HYDROXYACYLGLUTATHI ONE HYDROLASE [GLYOXALASE II) (GLX II) protein [Escherichia coli] egi[1786406 (AE000130)	5.7
;	• • • •	Xwnt-4) mRNA,			ydroxyacylglutathione	1
' I	U13183 c	omplete cds.	1.8		_	5.5

	Neare	st Neighbor (BlastN vs	. Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant Pr	Oteins)
SEC	- 1	DN DESCRIPTION	P VALUE			P VALU
├		No. (Cl() d				I VALU
116	S68944	Na+/Cl(-)-dependent neurotransmitter transporter	1.8	2276316	(Z96810) GLYT-1 LIKE [Homo	
		Rat calcium channel		2270310	sapiens]	5.5
		alpha-1 subunit (rbB	- [j	_	
117	1400005	I) mRNA, complete	1		(AF067607) Similar to cuticular	
11/	M92905	cds.	1.8	3165522	collagen; C18H7.3	5.5
118	X12429	Xenopus laevis U1 70K gene exon 10			(AF015685) reverse	<u> </u>
	1 112429	70K gene exon 10	1.8	2735957	transcriptase domain protein	3 .3
	ł .	•	1	. ,	cyclin A/CDK2-associatd	
	ł	1	ļ	·	protein P19 (RNA polymerase	
	I	1			elongation factor)	
			1		(SW:SKPI_HUMAN); cDNA	
	1				EST EMBL:T00114 comes	
	1		i l		from this gene; cDNA EST	
	}		1 1		yk390f11.5 comes from this	
	l	1	1 1		gene; cDNA EST yk402e11.5	
İ		1	1 1		co	
		1	1 1		>gi 3877216 gnl PID e1346850	
- 1		Mouse hepatitis virus] [protein P19 (RNA polymerase	
		genomic RNA for	1		elongation factor) gene; cDNA	
ĺ		spike protein, partial	1		EST yk390f11.5 comes from	
119	D83333	cds	1.8	207/570	this gene; cDNA EST	
			1.0	3876559	yk402e11.5 co	3.3
- 1		Cervus elaphus			(Z99942) similar to von	1
- 1		REDDEER	1		Willebrand factor type A domain; cDNA EST yk412d4.5	
1		mitochondrial D-	1		comes from this gene; cDNA	ľ
		loop, complete	ľ		EST yk412d4.3 comes from this	
20	AF016972	sequence	1.8	3878057	gene	٠, ١
- 1					HYPOTHETICAL 21.0 KD	3.2
ł		·	ł	•	PROTEIN IN RPS3-PSD1	
- 1			1		INTERGENIC REGION	1
ı		I	1		>gi 2132762 pir S63129	ı
]	j]		probable membrane protein	ł
	j		J		YNL174w - yeast	i
- [į,	Oncorhynchus mykiss	ſ		(Saccharomyces cerevisiae)	-
į		mRNA for rtSox24,	1	•	>gi 1302152 gnl PID e239548	- 1
1		complete cds	1.8	1720905	(Z71451) ORF YNL174w	
			- •••	1730805	[Saccharomyces cerevisiae] A-TYPE INCLUSION	2.5
1	1	Haemophilus	1			ĺ
- [ļi	nfluenzae Rd section	1	i	PROTEIN (ATI) camelpox virus >gi 62381 (X69774)	- 1
2		59 of 163 of the	j		84kDa A-type inclusion protein	
	U32844 c	omplete genome	1.8	728910	[unidentified]	1

<u></u>	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghhor (Bluery va Non B. L.	
SEC	2		T	1.04.036.1461	ghbor (BlastX vs. Non-Redundant P	roteins)
ID.	ACCESSIO	DN DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VAL
 	+					I VAL
ŀ	1	Human ionizing				
	1	radiation resistance	1	1		1
123	7718201	conferring protein			ribosomal protein YS7 homolog	İ
123	U18321	mRNA. complete cds	5. 1.8	2133273	Emericella nidulans	
	1	Human cystic fibrosis	_		·	I.4
	j	mRNA, encoding a	'	j]
	Į.	presumed			1	
	1	transmembrane	1	ł	1	i
		conductance regulator	<u>.</u>]			
	f	(CFTR). > ::	' }		[
		gb I11500 I11500	ľ			
		Sequence 1 from				
124	M28668	Patent US 5407796	1.8	22.125	filaggrin precursor - mouse	
			1.0	90492	(fragment)	0.87
					PROBABLE PROTEIN	
1	•			•	DISULFIDE ISOMERASE PS	
- 1		Mus musculus NSD1	ŀ		PRECURSOR >gi 1065461	
- 1		protein mRNA,			(U40411) Similar to protein	
125	AF064553	complete cds	1.8	2501207	disulfide-isomerase.	
				2301207	[Caenorhabditis elegans] REGULATORY PROTEIN	0.87
			1		BRLA (BRISTLE A PROTEIN)	
- 1			- 1		>gi 83718 pir A28913	
		Human mRNA for	· · · · · · · · · · · · · · · · · · ·		regulatory protein brlA -	
26	4 B0000 4 4	KIAA0316 gene,	- 1		Emericella nidulans >gi 168029	
20	AB002314	complete cds	1.8	115131	(M20631) brlA protein	0.84
- 1		Homo sapiens			(says and protein	0.84
		(subclone 10_d2 from	į.			
27	L42096	P1 H21) DNA sequence.			metalloproteinase 1 (EC 3.4.24	
		R.norvegicus renin	1.8	2135624) - human	0.65
28	M37278	gene, exons 1-9	, ,		(AF109907) S164 [Homo	
		Artificial sequences	1.8	4050087	sapiens]	0.58
- 1		DNA for ART 2	1		(L13442) cysteine-rich extensin-	
<u>.</u>	X82879	consensus	1.8	310020	like protein-4 [Nicotiana	
				310929	tabacum]	0.52
- 1	Į:	Homo sapiens mRNA	1			
	ļ.	for CRM1 protein,	1		(4 1010702) 34 54 5	
0		omplete cds	1.8	3559944	(AJ010792) Muc5AC protein	
1	7				[Mus musculus]	0.38
1		vlus musculus	i	1.	(AE000766) enolase-	
	S S	epiapterin reductase	ĺ		phosphatase E-1 (Aquifex	
	U78076 lo	ene. exons 1 and 2	1.8	11	huoshiarase E-T [Wdittex	

		Near	est Neighbor (BlastN v	/s. Genbank)		Nearest Na	ighbos (Dl	
S	EQ			1		. remest IACI	ghbor (BlastX vs. Non-Redundant I	Proteins)
	- 1	ACCESSI	ON DESCRIPTION	N P VAL	.UE	ACCESSION	DESCRIPTION	P VALUI
							COLLAGEN ALPHA I(VIII)	1
	- 1		1	- 1	1		CHAIN PRECURSOR	
	- 1		Paramecium 168G	.	- 1		ENDOTHELIAL	1
ł	- 1		gene for 168G	'			COLLAGEN)	ł
13	32	X52133	surface protein		- 1		>gi 105686 pir \$15435 collagen	. [
	7	100.55	surface protein	1.8		115316	alpha I(VIII) chain precursor - (U61944) coded for by C.	0.073
- 1	- 1			İ			(U61944) coded for by C.	
1	- 1			- 1	- 1		elegans cDNA ykl 12f3.5; codec	i]
- 1	- 1		1	İ			for by C. elegans cDNA	l
1				1	ł		cm21d2; coded for by C.	j
ı	- 1] '	I	- 1	1.	elegans cDNA CEESR07F;	1
	-			ł			coded for by C. elegans cDNA	
	1		Human desmoplaki		- 1		yk112f3.3; coded for by C.	
133	3	M77830	mRNA, complete co	. !	ı		elegans cDNA CEESR29F	
	\top		Plasmodium berghe	is. 1.8		1397246	[Caenorhabditis elegans]	le-04
134	1	<u>AJ2</u> 24150	EF-Lalpha A-gene	1.8	- 1	135084	(U43192) myosin II heavy chain	
	T		Taipila 71 gene	1.8	+	1353761	[Naegleria fowleri]	2e-05
1	1		Mus musculus	1	- 1		(7,500,10)	
j			somatostatin recepto	or l			(U58748) similar to potential	j
1	1		2 gene, exon1 and 5		- [transmembrane domains in S.	- 1
135	1_	AJ005518	flanking region	1.8		1326350	cerevisiae nulcear division	1
	1		Ralstonia eutropha	1	+	1320330	RFT1 protein (SP:P38206)	2e-08
1	1		megaplasmid pHG1	1	-	•	1	
1	1		nitric oxide reductase	•			1	1
١	1		(norB) gene,				(41 021121)	- 1
136	A	F002217	complete cds	1.8	1	3393018	(AL031174) hypothetical	
l	1		Caenorhabditis		7	0070010	protein	2e-08
137	١.		elegans cosmid	ı	1		(AL033388) 3-oxoacyl-[acyl-	
137	<u>A</u>	F039035	C53A3	1.8	1	3850109	carrier-protein]-synthase	
			S da		Т		proteing-synthase	3e-11
		i	S.domesticus	1	1		1	- 1
			immunoglobulin	1	1			. 1
			rearranged gamma	1			(AL022600) putative mannose-I	
138	N.		chain mRNA, VJC		1		phosphate gaunyl transferase	1
		-51/09	region, complete cds.	8.1	 	3080527	[Schizosaccharomyces nombel	3e-14
1					1		HYPOTHETICAL 24.2 KD	
j		1			[1.	PROTEIN C13A11.03 IN	1
139	_ Y	11106	P.pastoris PYC1 gene	1.0	l	J·	CHROMOSOME 1 >gil984224	
7			Human putative	1.8	 -	1175412	(751000)	le-15
ł			Ca2+/calmodulin-		1			
			lependent protein		l		(AL021687) putative protein	1
- 1		k	inase kinase gene, 3'		l		[Arabidopsis thaliana]	ł
			lanking region,			[>	ei 2832633 gnl PID e1249651	ł
140	U		artial sequence	1.8		2828280	AL021711) putative protein	
					0	2020200	Arabidopsis thaliana]	3e-17

1 ~-		rest Neighbor	(BlastN vs. (Genbank)	Nearest Neis	hbor (BlastX vs. Non-Redundant	
SE	· 1	ŀ				Vs. Non-Redundant	Proteins)
III	ACCESS	ION DES	CRIPTION	P VALU	ACCECCION.		1
		Plasmod		T VALUE	ACCESSION	DESCRIPTION	P VAL
		falciparı		 -			
	1	chromos		1			
i I	Ĭ		7 of 73 of				1
	1	the comp		1 .			
141	AE0014			1		(U95973) endomembrane	
	T	- Judgenee		1.8	1931647	- protein EMP70 precusor isolog	2e-20
	1	1		ĺ	1	IN VPOTHETICAL 75.5 KD	'
	1	- 1			1	PROTEIN C14A4.3 IN	1
	1	ł	í			CHROMOSOME II	
	1	Rat N-me	thyl-D-		j	>gi 3874230 gn1 PID e1351618	1
	i	aspartate			1 .	protein (Swiss Prot accession	1
	1	(NMDAR	1) gene		i .	number P38376); cDNA EST	1
142	L19708	first exon.	-, 50.10,	1.8		yk220e10.5 comes from this	
		Table Gradit		1.8	1731181	gene [Caenorhabditis elegans]	3e-21
•		1	1		ł	(Z81103) predicted using	1 21
	Î	Ī	Ī		Í	Genefinder, cDNA EST	1
	ļ	P.schwarz	i			yk303g11.5 comes from this	1
	1	mitochond	rial cyth			gene; cDNA EST yk303g11.3	İ
143	Y10728	gene, parti		1.8	22-21	comes from this gene	1
		, , , , , , , , , , , , , , , , , , ,	"	1.8	3878644	[Caenorhabditis elegans]	le-28
ł	İ	Homo sapi	ens mRNA				
- 1		for KIAA0	293 gene				l
44	AB006631	partial cds		1.8	4176600	(AL031177) dJ889M15.3 (novel	İ
- 1		Mus muscu	ılus I3	1.0	4176500	protein)	7e-45
ł		protein mR	NA.	I			
45	AF106967	complete co		1.7	-NO.177		
- 1		Archaeoglo	bus		<none></none>	<none></none>	<none></none>
1		fulgidus sec		- 1			
		172 of the c	omplete	1			
16	AE001073	genome		1.7	<none></none>		
		Pseudomon			- CHOINES	<none></none>	<none></none>
1		lemoignei p	oly(3-	1	ļ		
		hydroxybuty	rate) .	1		1	
- 1		depolymeras	e A	- 1		i	
1		precursor (pl	haZ5)	1	1	I	
1		gene, comple		j	1	ł	
1	ļ	and glycerol-	.3.		}		
1		phosphate-	1		į	1	
1	·	dehydrogena	se		Į	ľ	- 1
. I	1	homolog, cor	nplete	1	1	. 1	ľ
4	U12977	cds.	1	1.7	_ <none></none>		i
1		Mus musculu			NAONES	<none> <</none>	NONE>
1		SK/CamRk)		1			
	1.	germline IgK	chain	i	į.		- 1
	M27038	Sermine 18K	CHEER [J.	

25.50	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ			Citoank)	tremest ineign	OUT (Blasta vs. Non-Redundant l	roteins)
ID	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALU
		H.sapiens HBF-1				
1		mRNA for				
149	X74142	transcription factor Streptococcus	1.7	<none></none>	<none></none>	<none></none>
		thermophilus DeoD			1	' '
		gene, partial cds and				
		EpsA, EpsB, EpsC,				
		EpsD, EpsE, EpsF,				
		EpsG, EpsH, Epsl,				}
		EpsJ, EpsK, EpsL,				
li		EpsM, Orf14.9		,		· · ·
150	U40830	protein genes, complete cds	1.7	-NONTE		
150	U-0630	Rabbit lg germline	1.7	<none></none>	<none></none>	<none></none>
		gamma H-chain			1	
		(allotype d12,e15) C-			1	
151	L29172	region gene. 3' end.	1.7	<none></none>	<none></none>	<none></none>
152	M19045	Human lysozyme		2705		
132	14113(14)	mRNA, complete cds.	1.7	<none></none>	<none></none>	<none></none>
		Borrelia burgdorferi			1	1
ı		(section 45 of 70) of				
153	AE001159	the complete genome	1.7	<none></none>	<none></none>	<none></none>
Ī		Plasmid pFdA (from				
ł		Fremyella	j]
ļ		diplosiphon) DNA sequence, including	i			1
ŀ	1	unidentified cds and	1			
154		stem loop.	1.7	· <none></none>	<none></none>	<none></none>
					NI TOTAL	-HONE>
ı		Arabidopsis thaliana	1			
- 1		Columbia GTP	• •			
		binding protein beta	.	·		
155		subunit (AGB1) mRNA, complete cds.	1.7	ANONE	-11017	
	012232	incian, complete cus.	1./	<none></none>	<none></none>	<none></none>
1		Arabidopsis thaliana	İ			[]
		ATPK6 mRNA for	1			
1		ibosomal-protein S6				
,,,		inase homolog,				
156	D42056 c	omplete cds	1.7	<none></none>	<none></none>	<none></none>
	l _s	Rhizobium		İ		
	1	eguminosarum prsD,		1		
157		rsE, ORF3 genes	1.7	<none></none>	. <none></none>	<none></none>

	Nearest	Neighbor (BlastN vs. G	ienbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant l	Denosiu-1
SEQ					Tool (Diasot vs. Ivon-Redundant)	roteins)
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE
	 	 				
	1	Spinacia oleracea				
150		heat shock 70 protein				
158	AF039084	protein, complete cds	1.7	<none></none>	<none></none>	<none></none>
		R.norvegicus gene for				
150	7:065	catechol				1
159	Z12651	methyltransferase	1.7	<none></none>	<none></none>	<none></none>
		Fringilla coelebs		-		
		mitochondrial control				İ
160	A T0000 60	region, partial				
160	AF002968	sequence	1.7	<none></none>	<none></none>	<none></none>
		b				
		Borrelia burgdorferi	٠.		·	İ
161	47700111	(section 46 of 70) of				
161	AE001160	the complete genome	1.7	<none></none>	<none></none>	<none></none>
- 1		Methanococcus				
j		jannaschii section 95	ı		,	
162	1167663	of 150 of the				
102	U67553	complete genome	1.7	<none></none>	<none></none>	<none></none>
163	1496247	S.ruminantium				
103	M86247	plasmid pS23 DNA. oIL-8=interleukin-8	1.7	<none></none>	<none></none>	<none></none>
			ŀ			
164	S74436	[sheep, spleen cells,				
	374430	mRNA, 1435 nt] Candida maltosa	1.7	<none></none>	<none></none>	<none></none>
		ALK7 (CYP52A10)	- 1			
ľ		and ALK8 complete	- 1			1 1
165		eds				1 1
		Geotrichum	1.7	<none></none>	<none></none>	<none></none>
		candidum NRRL Y-				
- 1		553 lipase gene,	I		220	
166		partial cds.	1.7	201046	230k bullous pemphigoid	
			1./	321245	antigen BPM1 - mouse	9.3
	·],	H.sapiens CpG DNA	j			
- 1		clone 114a4, reverse			(1/66209) hamage	
167		read cpg 14a4.rt la .	1.7	1854675	(U66298) bone morphogenetic	
<u> </u>		ордилатици.		10340/3	protein-6 [Rattus norvegicus]	9.1
	ĺ		1		PROTEIN BAT2 MHC class III	
i	1	j	1		histocompatibility antigen HLA-	i
	ĺ		- 1		B-associated transcript 2 -	
1	1		1		human >gi 179339 (M33509)	
	[1		HLA-B-associated transcript 2	
	A	Agrobacterium	J		(BAT2) [Homo sapiens]]
- 1		imefaciens conjugal	ı		>gi 179345 (M33518) HLA-B-	
- 1		ansfer region 1	1		associated transcript 2 (BAT2)	
68		enes	1.7	1352066	[Homo.sapiens]	, I
				1302000	[110000.Supiciis]	9.1

	Neare	st Neighbor (BlastN vs.	Genhank)	Nearest Mai-	ht. and	
SE			John Mark	Incarest Neig	hbor (BlastX vs. Non-Redundant P	roteins)
ID	-	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-					PROTOPORPHYRINOGEN	I VALUE
	1				OXIDASE (PPO) yeast	
- 1	1	1	ł	ľ	(Saccharomyces cerevisiae)	
1		Comment		j	>gi 603606 (U18778) Hem14p:	1
ł	İ	Caenorhabditis	Ţ		protoporphyrinogen oxidase	1
	į.	elegans cosmid	1		[Saccharomyces cerevisiae]	l
ı	1	Y12A6A, complete			>gi 1403536 gn1 PID e249333	i i
ł	Į.	sequence [Caenorhabditis			(Z71381) protoporphyrinogen	1
169	AL023827		1		oxidase [Saccharomyces	
1	162023827	X.laevis mRNA for	1.7	731440	cerevisiae]	8.9
1		glutathione	i I			
1	1	synthetase, large				
170	X69662	subunit	1.7	40.000.00	(AC005897) hypothetical	j
		S.cerevisiae	1./	4038057	protein [Arabidopsis thaliana]	8.8
1	ì	chromosome II				
1		reading frame ORF			(115515)	1
171	Z35824	YBL063w	1.7	3021450	(Y15515) prdl-a [Hydra	1
				3021430	vulgaris] COLLAGEN ALPHA I(XII)	7.0
	1		ı		CHAIN PRECURSOR I(XII)	1
1	ĺ		i		chain - chicken	•
1	ŀ		i		>gi 222811 gnl PID d1001160	
		Cowpea chlorotic	j		gallus)	j
1 1		mottle virus (CCMV)			>gi 2326442 gn1 PID e39435	l l
172	1465100	la protein gene.			(X61024) collagen type XII	1
1/2	M65139	complete cds.	1.7	2506307	alpha I chain [Gallus gallus]	7.0
		ł	i		HYPOTHETICAL 10.0 KD	
		Ì	[PROTEIN IN ALPA-GABD .	· 1
		Drosophila distal BX-	İ	j	INTERGENIC REGION (F87)	- 1
		C region (bithorax	•		>gi 1033124 (U36840)	i
J		complex) pH189 5	į		ORF_f87 [Escherichia coli]	
173	77.00	region;	, ,		>gi 1788982 (AE000348) orf,	1
	-210000	region,	1.7	1723625	hypothetical protein	6.9

ACCESS	rest Neighbor (BlastN vs.	T		Cigipor (BlastX vs. Non-Redundant l	roteins)	
- 1	IONI DESCRIPTION			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
	ION DESCRIPTION	P VALU			P VALU	
		+			Ť	
1			1	(Z46792) similar to lethal(1)		
1	- 1	İ	1	discs large-1 tumor suppressor	1	
1	Ī	1	1	protein-like repeats; cDNA EST	:	
ł	1	1	1	EMBL D33495 comes from this		
i	ļ	1	1	gene; cDNA EST	ł	
l	1	1	ı	EMBL:D35117 comes from this	I	
	1	ł		gene; cDNA EST	ĺ	
	1	i	1	EMBL:D36356 comes from this	1	
	· ·	1	1.	gene; cDNA EST EMB	1	
			j	>gi 3879984 gnl PID e1351767]	
				suppressor protein-like repeats:	-	
		ŀ	1	cDNA EST EMBL:D33495		
			1	comes from this gene; cDNA		
	S.cerevisiae		1	EST EMBL:D35117 comes		
	1]	from this gene; cDNA EST		
Z46255	1	1.7	20222	EMBL:D36356 comes from this		
			38/5228	gene; cDNA EST EMB	6.7	
				THYMIDINE KINASE		
U01066		17	125440	saimiriine herpesvirus 1 (strain		
			123448	11[Onc]) >gi[60341	6.7	
				1		
T 72 4- 42	homeobox protein			(II38184) ATPage subverse		
U34743	mRNA, complete cds	1.7	1022918	(Trypanosoma cruzi)		
				(2.) panosonia ciuzij	6.7	
		•		(AL023862) hypothetical	j	
1114662		1		protein SC3F9.07 (Strentomyces		
V 1 7002	Homo sapiere	1.7	3218378	[coelicolor]	6.7	
		1				
AB017006				(U64859) glutamine-rich protein	j	
	pareiai eus	1./	1465855	[Caenorhabditis elegans]	6.7	
	Brassica oleracea var	- 1				
		- 1		DV0	I	
	intrinsic protein	ł		DYNEIN HEAVY CHAIN,	į	
	bobTIP26-1 mRNA.			C T I OSOLIC (DYHC) dynein	J	
U92651	complete cds	1.7	3023675			
			3423013	[Schizosaccharomyces pombe]	6.6	
	Lytechinus varicgatus	[(M58520) and 1.4 bis		
F000634	mRNA, complete cds	_1.7	148574		6.6	
	U01066 U34743 U14662 AB017006	Human CD4 promoter, partial sequence Phalaenopsis sp. 'hybrid SM9108' homeobox protein mRNA, complete cds Baboon herpesvirus HVP2 gB glycoprotein (UL27) gene, complete cds. Homo sapiens PMS2L15 mRNA, partial cds Brassica oleracea var. botrytis tonoplast intrinsic protein bobTIP26-1 mRNA, complete cds Lytechinus varicgatus notch homolog	Chromosome VI Ilambda clone. Human CD4 promoter, partial sequence. 1.7 Phalaenopsis sp. 'hybrid SM9108' homeobox protein mRNA, complete cds HVP2 gB glycoprotein (UL27) gene, complete cds. Homo sapiens PMS2L15 mRNA, partial cds Phassica oleracea var. botrytis tonoplast intrinsic protein bobTIP26-1 mRNA, complete cds 1.7 Lytechinus varicgatus notch homolog	chromosome VI lambda clone. Human CD4 promoter, partial sequence. 1.7 125448 Phalaenopsis sp. 'hybrid SM9108' homeobox protein mRNA, complete cds HVP2 gB glycoprotein (UL27) gene. complete cds. Homo sapiens PMS2L15 mRNA, partial cds. Brassica oleracea var. botrytis tonoplast intrinsic protein bobTIP26-1 mRNA. U92651 Lytechinus varicgatus notch homolog	EMBL:D33495 comes from this gene; cDNA EST EMBL:D3636 comes from this gene; cDNA EST EMBL:D3636 comes from this gene; cDNA EST EMBL:D3636 comes from this gene; cDNA EST EMBL:D3636 comes from this gene; cDNA EST EMBL:D36317 comes from this gene; cDNA EST EMBL:D36317 comes from this gene; cDNA EST EMBL:D36356 comes from this gene; cDNA EST EMBL:D363517 comes from this gene; cDNA EST EMBL:D363517 comes from this gene; cDNA EST EMBL:D363517 comes from this gene; cDNA EST EMBL:D363517 comes from this gene; cDNA EST EMBL:D363517 comes from this gene; cDNA EST EMBL:D363517 comes from this gene; cDNA EST EMBL:D363517 comes from this gene; cDNA EST EMBL:D363517 comes from this gene; cDNA EST EMBL:D36351 for the partial das das das das das das das das das das	

	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant P	
SE					Elloof (Blasta Vs. 140h-Redundant P	roteins)
11	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-	+	A -a bird i - d - 10				1. ************************************
ł		Arabidopsis thaliana				
- 1		anthranilate synthase				ł
18	1 M92354	alpha subunit gene, complete cds.	1	1	blue light photoreceptor	ľ
	14192554	Hordeum vulgare	1.7	738308	[Arabidopsis thaliana]	6.5
1	1	genomic DNA	1	i	(AC002411) Strong similarity to	
1	1	fragment; clone	l	ľ	myosin heavy chain gb/Z34293	ĺ
1 18:	AJ234856		1		from A. thaliana. [Arabidopsis]
	1205 1030	Stercorarius	1.7	3142302	thaliana)	6.5
- 1	1	parasiticus bird J33	!			
ı		cytochrome b protein,			21122	
183	U76827	partial cds	1.7	2412010	(Y17034) Bassoon [Mus	
		Saccharomyces	1.7.	3413810	musculus)	5.4
1	Ī	cerevisiae Ttp1p			İ	
1	1	(TTP1) gene,			(T 24402) II	
184	U05211	complete cds.	1.7	403173	(L24492) lipoprotein	
1				103173	[Rhodococcus erythropolis]	4.9
1	1	Homo sapiens	1		ENDOGLUCANASE TYPE K	
1	1	TRRAP protein			PRECURSOR (ENDO-1,4-	}
1.05	1	(TRRAP) mRNA,			BETA-GLUCANASE)	
185	AF076974	complete cds	1.7	1170140	(CELLULASE)	4.1
	1	1				7.1
1	Į	1	I		DNA ADENINE METHYLASE	
1	j	Aquifex aeolicus			site-specific DNA-	
ĺ		section 85 of 109 of			methyltransferase (adenine-	
186	AE000753	the complete genome			specific) dam methylase gene	- 1
	12200733	the complete genome	1.7	1169357	product [Vibrio cholerae]	4.0
	- ,	Tupaia glis	1			
		apolipoprotein AI	J			
		prepropeptide	J			
. 187		mRNA, complete cds	1.7	3355682	(AL031124) putative secreted	1
				3333082	lyase	4.0
		Human germline IgK	- 1			J
		chain gene V3-region,	•		(AB004534) pi003	
188	M23090	clone Humkv328h5	1.7	2257483	[Schizosaccharomyces pombe]	40
ı	j				(Semzosacenaromyces pomoe)	4.0
1	J,	\d?=1.	l		myotonic dystrophy kinase -	
- 1		Mink enteritis virus	1		mouse (fragment) kinase, DM-	1
		intigenic type 2	j		kinase {C-terminal, alternatively	1
i		apsid protein genes	j		spliced, clone delta II,III,IV,V}	.]
189		VP1 and VP2,	l		[mice, brain, Peptide Partial,	1
**/	M24001 c	omplete cds.	1.7	2143504	474 aa] [Mus sp.]	3.9
190		I.sapiens CST4 gene or Cystatin D			(U37273) winged helix protein	
	1.02204 11	or Cystatiii D	1.7	1766075	CWH-2 [Gallus gallus]	3.1

	Nearest	Neighbor (BlastN vs. (Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant		<u> </u>	
SEQ			T	carest iver	Hibbi (Blasta Vs. Ivon-Redundant P	rateins)	
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
-	 				HYPOTHETICAL 11.7 KD	I VALUE	
		P.falciparum complete gene map o plastid-like DNA (IR	f	-	PROTEIN C6B12.13 IN CHROMOSOME I		
191	X95276	B) Rat PMSG-induced	1.7	3219951	>gi 2330843 gnt PID e334047 pombe]	3.0	
192	D84487	ovarian mRNA, 3'sequence, N10	1.7	173164	(J02719) valy!-tRNA synthetase [Saccharomyces cerevisiae]	2.3	
193	L14851	Rattus norvegicus neurexin III-alpha gene, complete cds.	I.7	3323586	(AF060869) single-strand binding protein [Salmonella typhimurium]		
		Xenopus laevis/gilli hybrid pseudo-IgH			· ·	2.3	
194	M97002	chain gene, V region, clone LG7G342A. Bactinus unuringiensis	1.7	2118407	MHC sex-limited protein - mouse (fragment) musculus]	2.3	
		delta-endotoxin (CryA(a)) gene, 5' end. > :: gb I34520 I34520 Sequence ! from patent US 5596071 > :: gb I39790 I39790 Sequence ! from patent US 5616495 > :: gb AR008487 AR008			HYPOTHETICAL 53.4 KD PROTEIN D1054.13 IN		
195	L07025	487 Sequence 1 from patent US 5753492	1.7	2496940	CHROMOSOME V >gi 3875316 gn PID e1344967	1.8	
196	. (nsulin-like growth actor II {intron 7} human, Genomic, 702 nt]	1.7	3327038	(AB014512) KIAA0612 protein [Homo sapiens]		
	D	luman (lambda) NA for			Fv Fragment (Murine Se155-4) Complex With The Trisaccharide: Alpha-D- Galactose(1-2)[alpha-D- Abequose(1-3)]alpha- D- Mannose (P1-Ome) (Part Of	1.8	
197		nmunoglobulin light nain	1.7	ľ	The Cell-Surface Carbohydrate Of Pathogenic Salmonella)	1.8	

-	Nearest	Neighbor (BlastN vs. G	ienbank)	Nearest Neigh	ibor (BlastX vs. Non-Redundant Pr	manine'
SEO		Trengmoor (Diagna Vis. C	I I	ricarest ricigi	Blastic Vs. Non-Redundant Pi	oteins)
ID ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	
-	1.1000001011	Plasmid pFdA (from	I VALUE	ACCESSION	DESCRIPTION	P VALUE
1-	 	Fremyella				<u> </u>
ł		diplosiphon) DNA				
ı	1	1 -				l
İ		sequence, including unidentified cds and		·		l
198	L17027	1		1000700	poliovirus receptor-related	ł
170	L17027	stem loop. Caenorhabditis	1.7	1082702	protein - human	1.4
		elegans cosmid				
]	H22D14, complete				
]	İ	sequence			(AE060442)	ŀ
		[Caenorhabditis			(AF069442) putative inhibitor	
199	AL022273	[elegans]	1.7	3004606	of apoptosis [Arabidopsis	
	ALUZZZIS	ciegansj	1./	3924605	thaliana}	1.4
i i		Drosophila	• •			
		melanogaster cut			(707242) DNIA 1: 4:	
200	U89926	gene, partial sequence	1.7	2245100	(Z97343) DNA-binding protein	
200	007720	gene, partial sequence	1./	2243100	homolog PROTEIN KINASE C	1.3
					SUBSTRATE, 60.1 KD	
			ľ		PROTEIN, HEAVY CHAIN	
		H.sapiens gene for			(PKCSH) (80K-H PROTEIN)	
201	Z25749	ribosomal protein S7	1.7	2493459		
	223747	Fundulus heteroclitus		2493439	>gi 1215746 (AF048977) Ser/Arg-related	1.1
		lactate dehydrogenase			nuclear matrix protein [Homo	
202	U59841	B	1.7	3005587	sapiens]	0.82
		Rabbit mRNA for		3003307	3apicii3	0.02
		smooth muscle	1			
ł		calcium channel	1			
- 1		blocker (CaCB)	1		(AF082302) arabinogalactan-	
203		receptor	1.7	3883128	protein [Arabidopsis thaliana]	0.82
		Caenorhabditis			protein (r z z z z pos z z z z z z z z z z z z z z z z z z z	
- 1		elegans cosmid ·	1			1
	Į	C18B12A, complete			1	i
ľ	4	sequence	1			
		[Caenorhabditis	1	•	(D10123) core [Hepatitis C	·
204		elegans]	1.7	940397	virus]	0.80
Ī		Human XIST gene,				
		poly purine-	ł		(AF005370) large tegument	ļ
. 1		pyrimidine repeat	ļ		protein [Alcelaphine herpesvirus	ľ
205		region	1.7	2338027	1]	0.59
		Ovis aries beta-	I		(X65335) lacZ gene product	
206		actoglobulin gene	1.7	987050	[unidentified cloning vector]	0.45
		Homo sapiens	T	· · · · · · · · · · · · · · · · · · ·		
	The state of the s	roponin T (TNNT2)	į		(AF053947) unknown [Yersinia	
207	AF004419	gene, exon 13	1.7	2996364	pestis] >gi 3883090	0.22
	1.		- 1		TRANSDUCIN-LIKE	- 7
ſ		Gallus domesticus	j		ENHANCER PROTEIN I	1
200		DNA microsatellite	1		enhancer-of-split homolog TLE-	Ì
208	L43643 r	narker MCW119	1.7	464896	1 - human >gi 307510	0.20

i	Neare	est Neighbor (BlastN vs	Carbarth			
5	EQ	STITE ENDOT (BIASUT VS	. Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant	Proteins)
	D ACCESSIO	ON DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
\vdash						I VALUE
		S cerevisiae chromosome XII reading frame ORF			HYPOTHETICAL 123.9 KD PROTEIN C30D11.04C IN CHROMOSOME I >gi 2130411 pir S62562 hypothetical protein SPAC30D11.4c - fission yeast	
20	9 273278	YLR106c	1.7	1351657	nuclear pore complex protein	1
21	0 M22345	Mouse endogenous provirus gag, pol, an env region DNA. Escherichia coli K-1: MG 1655 section 250	d 1.7	2444455	[Schizosaccharomyces pombe] (AF020765) hypothetical protein [Myxococcus xanthus]	0.20
21	1 AE000360	of 400 of the complete genome	1.7	2736361	(AF039038) No definition line found [Caenorhabditis elegans]	0.12
212	2 AB020692	Homo sapiens mRNA for KIAA0885 protein, complete cds testis-determining	1	2605924	(AF029726) histidine kinase C [Dictyostelium discoideum]	0.094
213	S69429	gene/SRY homolog [Sminthopsis macroura=striped- faced dunnarts, Genomic, 855 nt] testis-determining	1.7	2499016	TONB PROTEIN >gi 1666536 (U23764) TonB [Pseudomonas aeruginosa]	0.092
214	S69429	gene/SRY homolog [Sminthopsis macroura=striped- faced dunnarts, Genomic, 855 ht]	1.7	2499016	TONB PROTEIN >gi 1666536 (U23764) TonB [Pseudomonas aeruginosa]	0.088
215	U67205	Mus musculus ACF7 neural isoform 3 (mACF7) mRNA, partial cds	1.7	2047349	(AF000198) weak similarity to HSP90 [Caenorhabditis elegans]	0.052
216	X98188	Artificial DNA sequence for mammalian lambda- neo minichromosome, 1400 bp	1.7	2493779	PUTATIVE CUTICLE COLLAGEN C09G5.6 collagen; cDNA EST yk244c3.5 comes from this gene; cDNA EST yk244c3.3 comes from this gene [Caenorhabditis elegans]	
217		Mus musculus putative CCR4 protein mRNA, partial cds	1.7		(U95973) hypothetical protein [Arabidopsis thaliana]	0.042

	Near	est Neighbor (Blast)	vs. Genha	nk)	Nearest Na		
S	EO		3. 3004	,	IAETIESI IASI	ghbor (BlastX vs. Non-Redundant E	Proteins)
- 1	D ACCESSI	ON DESCRIPTION	ON P	/ALUE	ACCESSION	1	
-							P VALUE
- [Homo sapiens a	lpha-			(AF045246) universal minicircl	
	ł	l type V collage	n		l	sequence binding protein	el
21	1 20000	(COL5A1) gene	. 5'			minicircle sequence binding	1
 	L38808	flank and exon 1		1.7	2895760	protein [Crithidia fasciculata]	1
i	İ					Pertandia fasciculata)	. 0.039
21	277151	B.napus mRNA		I		(K02576) salivary proline-rich	J i
1	9 Z 72151	AMP-binding pro	otein	1.7	190475	protein 1 [Homo sapiens]	0011
1	ŀ	n · -		7		2 (120mo sapiens)	0.011
1	1	R.norvegicus mR	NA	- 1		synapsin.IIb - human	1 1
220	NO.	for cysteine sulfii	nate	1		>gi 1594277 (U40215) synapsin	1 1
1 22	0 X94152	decarboxylase	1	.7	2136212	IIb [Homo sapiens]	
22	1 1 20055	Mouse stathmin g		. 1		(U97553) unknown [murine	0.008
-22	L20255	sequence.	1	.7 [*]	2317934	herpesvirus 68]	0000
1	1					7-3-125001	0.006
1	i	Rattus norvegicus		- 1		(U23175) similar to anion	<u> </u>
222	1 12600	glycine transporte	r	[exchange protein	i
1 222	L13600	mRNA, complete	cds. 1	.7	726403	[Caenorhabditis elegans]	0.000
223	A 1724150	Plasmodium bergi	iei			(U95094) XL-INCENP	0.003
-223	AJ224150	EF-1alpha A-gene	1.	7	2072290	[Xenopus laevis]	0.00
ł	i	butyrophilin [mice				(second mevia)	0.001
	1	lactating mammar	′ I	- 1		ĺ	- 1
224	530640	gland, mRNA Part	ial,	- 1		(AJ223010) Pmt2	1
-224	S80642	3193 nt]	1	7	2695746	[Schizosaccharomyces pombe]	0-04
		C.elegans unc-86	1	$-\Gamma$		(Composite of the Composition of	9e-04
	i i	gene encoding two	- 1	1		1	1
225	Maaaca	alternative proteins	•	i		(AB002369) KIAA0371 [Homo	
225	M22363	complete cds.	1.7		2224683	sapiens]	le-04
						(Z49909) similar to Prokaryotic	1e-04
226	X92123	M.musculus egt ger				ribonuclease PH	1
	A92123	exon l	1.7	L	3874232	[Caenorhabditis elegans]	3e-05
		Inomes - 1 Drs. o	j	ı		- Carlo (Inc.)	36-03
		Ipomoea nil PKn2		j		1	1
227	AB016000	(knotted-like gene)		-		(AF000422) TTF-I interacting	
=	AB010000	mRNA, complete co	s 1.7		2183083	peptide 5 [Homo saniens]	le-05
ſ			1	- 1		peptide 5 [Homo sapiens] (AL032643) similar to	16.03
- 1				- 1	į,	Uncharacterized protein family	1
- 1	ľ		1	- 1	[1	UPF0034, Double-stranded	1
ı	į		1	1	ļi	RNA binding motif; cDNA EST	ŀ
- 1	1		Ī	1	ls ls	k489b3.5 comes from this	ŀ
	l,		1	- 1	وا	ene; cDNA EST yk439g7.5	}
J		Course on DAIA f	.		10		
228	D 4 4 4 4 4 - 1	Bovine mRNA for ynaptocanalin I	1.7	Í	اد	omes from this gene	

	Neares	t Neighbor (BlastN vs.	Genbarik)	Nearest Neig	hbor (BlastX vs. Non-Redundant F	Protein \
SEQ					Ton-Accountant F	roteins)
ID	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	İ
			T. VILLOE	Necession	DESCRIPTION	P VALU
	i					
	1	Mus musculus TAFO	3 -			
	l .	1-like neuronal	1	ļ		
	1	glycoprotein (PCS)	İ		(AL022299) hypothetical	1
229	L01991	mRNA, complete cds	1.7	3006139	protein	4e-07
]	Tomato yellow leaf				46-07
		curl virus Thailand	1			1
ľ		isolate complete	1			1 .
220		genome (TYLCV-TH	[(AC005395) hypothetical	1
230	X63016	B-DNA)	1.7	3643608	protein [Arabidopsis thaliana]	le-07
		H.sapiens		·.		
		microsatellite repeat.	1			İ
		> ::	٠,		{	
		gb G34562 G34562				
		human STS SHGC-			extensin precursor (clone Tom L	
231	Z22802	51834	1.7	100210	4) - tomato esculentum]	4e-09
ľ		Human complement	1			
1		component C3	i 1			
		mRNA, alpha and	•			
232	V00765	beta subunits,	1		(AE000773) acetoin utilization	
232	K02765	complete cds.	1.7	2984320	protein [Aquifex aeolicus]	le-09
					(Z73102) predicted using	
- 1			Í	•	Genefinder; Similarity to	
ı					Bacillus subtilis DNAJ protein	
- 1		S.cerevisiae	Į.		gene; cDNA EST	
J		chromosome XV			EMBL:C12520 comes from this	
		reading frame ORF			gene; cDNA EST	
233	_	YOL076w		2072700	EMBL:D71409 comes from this	
\neg	2	102070W	1.7	3873700	ge	7e-11
	Į,	Pig mRNA for thimet	ſ		(VIESIX) Parada	
234		oligopeptidase	1.7	2622000	(Y15513) Prodos protein	
\neg			1./	2632098	[Drosophila melanogaster] (Z81070) cDNA EST	8e-13
	ſ	·			EMBL:C12730 comes from this	
	i	1			1	
	ľ		1		gene; cDNA EST yk200b6.5 comes from this gene; cDNA	·
	lo	Gallus gallus gene			EST yk349a12.5 comes from	l
		ncoding neurofascin.	j.		this gene [Caenorhabditis	
25		xons 9,10,11 & 12	1.7	3876421	elegans]	1
35		cerevisiae		2070421	cicani	3e-14
"	ĮS	-cerevisiae				
33		hromosome XVI			(U64605) C05D9 6 gaps	Į
36	c		·		(U64605) C05D9.6 gene product {Caenorhabditis	ł

1	Nearest 1	Neighbor (BlastN vs. G	enbank)	Nearest Neigh	ibor (BlastX vs. Non-Redundant P	mteins)
SEQ		Langer (Diagram of			- Con (Diasor vs. 14011-1404) [Call P	(Otenis)
ID ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE
					OLIGOSACCHARYL	
					TRANSFERASE STT3	
]]		Homo sapiens			SUBUNIT HOMOLOG	1
		genomic DNA, 21q			>gi 529357 (U13019) No	
		region, clone:			definition line found	1
237	AG000518	T171N23	1.7	1174468	[Caenorhabditts elegans]	6e-18
		Human mRNA for N-				
		acetylglucosaminyltra				ŀ
		nsferase V, complete			(D63877) KIAA0157 gene	
238	D17716	cds	1.7	961446	product is novel.	5e-19
		Cheilodactylus			1	1.
1 1		vittatus country USA:		,		
i I		Midway Island	٠.			
		cytochrome c oxidase				
		subunit I gene,				1
		mitochondrial gene			(17700 to 1000 to]
l i		encoding mitochondrial			(U70848) C43G2.1 gene	1
239	AE102512			1500055	product [Caenorhabditis	1
239		protein, partial cds Rattus norvegicus	1.7	1572756	elegans]	5e-40
		liver-specific				
		transporter gene,			(41 022228) 4110421/10 4	
240		promoter region.	1.7	4176443	(AL022238) dJ1042K10.4 (novel protein)	3e-49
	ESOTO;	promoter region.	<u>^:/</u>	4170443	(nover protein)	36-49
		H.sapiens mRNA for]
		Na-Cl electroneutral				
	1	thiazide-sensitive			(AC005546) R29425_1 [Homo	
241	X91220	cotransporter	1.7	3478637	sapiens]	6e-54
	ł		ľ		1	ł i
- 1]:	Rattus norvegicus				[
		calcium-independent			1 .	j
- 1		phospholipase A2	ŀ	•	· _	
242		mRNA, complete cds	1.6	<none></none>	<none></none>	<none></none>
1		Pea secd-borne				
1		mosaic virus RNA for	1		1	
		coat protein and		•		
243		polymerase (partial)	1.6	<none></none>	<none></none>	<none></none>
- 1		Rat leukocyte	1			
- 1		common antigen (L-	1			!
244		CA) gene, exons 1				,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
244		hrough 5.	1.6	<none></none>	<none></none>	<none></none>
i		Yeast (S.pombe)				
		dc25+ gene (mitosis	1			
245		nitiation), complete	,	101 m		
245	M13158 c	ds.	1.6	<none></none>	<none></none>	<none></none>

	Neares	Neighbor (BlastN vs. C	ienbank)	Nearest Neir	hbor (BlastX vs. Non-Redundant P	
SEC			1		to the section of the	roteins)
Ð	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALU
<u> </u>		Mycoplasma				1
Ì	1	genitalium section 34				
1		of 51 of the complete		•		
246	U39712	genome	1.6	<none></none>	<none></none>	<none></none>
i]				(Z81499) predicted using	1
1		Mouse Murine			Genefinder; cDNA EST	ŀ
1	i	urokinase-type			yk410e3.3 comes from this	.]
ļ	ł	plasminogen activator			gene; cDNA EST yk410e3.5	1
247	1415000	protein gene,			comes from this gene	
247	M17922	complete cds.	1.6	3875750	[Caenorhabditis elegans]	8.0
248	M00004	Human polymorphic			(Z84724) psd [Mycobacterium	
240	M89986	loci in Xq28.	1.6	3261710	tuberculosis]	6.4
249	M00004	Human polymorphic			inositol-polyphosphate 4-	
249	M89986	loci in Xq28.	1.6	2143805	phosphatase - rat	6.2
		Rattus norvegicus	ł			
250	U68725	Deleted in colorectal Cancer			(U51449) RING3 protein	
230	008723	Cancer	1.6	1256804	[Xenopus laevis]	5.8
			1		MALÉYLACETATE	
		P.platessa GSTA,	1		REDUCTASE Pseudomonas	
		GSTA1, GSTA2, and	1		cepacia >gi 643636 (U19883)	
251	X95199	PPTN genes	1.6	2015112	maleylacetate reductase	
		11 Th genes	1.0	3915113	[Burkholderia cepacia] HYPOTHETICAL 91 KD	4.9
			1		PROTEIN IN COB INTRON	
	i		1			
					>gi 2654230 gnl PID e1192341 (X02819) unidentified reading	
		D.melanogaster			frame [Schizosaccharomyces	
252	Y09103	RPA1 gene	1.6	3916021	pombel	
		T.aestivum		3310021	politice	4.8
- 1		mitochondrion fMet,	- 1			
		18S, 5S repeat unit			DYSTROPHIN-RELATED	
253	Z14078	DNA	1.6	2501668	PROTEIN 2 sapiens]	3.6
					2 NOTELLY 2 Supicing	3.0
	į	·	į		REPETITIVE PROLINE-RICH	
ĺ	1		1		CELL WALL PROTEIN I	l
ſ	ľ		· 1		PRECURSOR	
- 1		i	1		>gi 81809 pir A29324 proline-	f
j	f	luman mRNA for	į		rich protein precursor - soybean	f
۱ ،		CIAA0316 gene,	1		>gi 170049 (J02746) proline-	ł
254	AB002314 c	complete cds	1.6	130997	rich protein [Glycine max]	2.8
	[j		ENV POLYPROTEIN	
	1.		J		PRECURSOR (COAT	
		luman muscle	1		POLYPROTEIN) [CONTAINS:	Ī
255		reatine kinase gene			COAT PROTEIN GP62: COAT	i
	14171499	CKMM), exon 2.	1.6	119399	PROTEIN GP40]	2.2

	Nearest	Neighbor (BlastN vs. C	Genbank)	Negrest Nair	yhhor (PlantV us N - P	
SEC			T T	ivernest ideli	ghbor (BlastX vs. Non-Redundant	Proteins)
ID	ACCESSION	N DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALU
-	 	Po-dia burda 6				
İ	1	Borrelia burgdorferi	1			
256	AE001164	(section 50 of 70) of the complete genome		105000	(AF109907) hypothetical	1
	1.0001104	The complete genome	1.6	4050089	protein [Homo sapiens]	1.5
	1	M.musculus				
l	j	rearranged T-cell	ŀ			
]	receptor beta variable			(AF080090) semaphorin IV	ł
257	X61757	region (Vb17a)	1.6	3377766	isoform b [Mus musculus]	1.2
		T.cruzi tandemly			(istas mascaras)	1.2
		repeated gene		,		1
	1	encoding an 85 kDa		,	(AF043695) similar to zinc	1
	İ	antigen with homology to heat	٠.		metalloprotease family of	1
258	M15346	shock proteins.			peptidases [Caenorhabditis	1
	14115540	snock proteins.	1.6	2804437	elegans]	0.41
		Rattus norvegicus				
		sodium channel	j			
		protein 6 (SCP6)	I		(AF018081) type XVIII	1
259	L39018	mRNA, complete cds	1.6	2920535	collagen [Homo sapiens]	0.037
						0.037
		Human leukocyte	i		ł	į i
Į		adhesion protein	- 1			j
260	M29483	p150,95 alpha subunit gene, exons 7 - 15.	1	*****	(U49082) transporter protein	
		gene, exons / - 15.	1.6	1840045	[Homo sapiens]	2e-09
[Aspergillus niger beta	i		·	[[
ł	1	D-fructofuranosidase	i]]
		(suc1) gene, one	- 1		(AF071527) putative calcium	
261	L06844	exon.	1.6	4206210	channel [Arabidopsis thaliana]	9e-10
- 1	1	a				70.0
1		Chicken aldolase B	i i		(AF040640) similar to peptidase	I
1		gene, complete cds, clones lambda-	- 1	•	family C19 (ubiquitin carboxyl-	
262	3 4 4 4 4 4 4	C(11.1.4).	1.6	2746776	terminal hydrolase)	1
		Human gene PRB3L		2746775	[Caenorhabditis elegans]	1e-31
- 1		or proline-rich				
263	X07881 F	protein G1	1.5	<none></none>	<none></none>	<none></none>
	T.	0				VI ONE
		Vicotiana tabacum	1	•	(Z99271) similar to Reverse	ĺ
264		JMP synthase (pyr5-			transcriptase comes from this	[
-) mRNA, partial eds	1.5	3880923	gene [Caenorhabditis elegans]	0.50
		uclear protein	J			
		ilP45 mRNA.			(US0761) COLET	i
65		omplete cds	1.4		(U58761) C01F1.6 gene product	
		<u></u> -		1330394	[Caenorhabditis elegans]	8.9

	Nearest	Neighbor (BlastN vs.	Canbank)	NonN		
SEQ		Treignor (Diastr Vs.	Genbankij	Mearest Neig	hbor (BlastX vs. Non-Redundant F	roteins)
,						ŀ
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
<u> </u>					POTASSIUM-	1
					TRANSPORTING ATPASE	
	l			j	BETA CHAIN (PROTON	ļ
1			İ	ľ	PUMP) (GASTRIC H+/K+	1 .
1	1		İ	1	ATPASE BETA SUBUNIT)	1
1	j	Rice tungro	ł	ŀ	3.6.1.36) beta chain - human	1
1		bacilliform virus	ł	ł	>gi 184105 (M75110) H,K-	İ
		Serdang strain,			ATPase beta subunit [Homo	
266	AF076470	complete genome	1.4	1703461	sapiens)	8.9
		C.jacchus interferon				
1 2/2		gene for interferon	1		(U28832) US 10 [Gallid	1
267	X64659	gamma	1.4	1486485	herpesvirus 1] >gi 1486497	6.8
1 1		Schistosoma	1			
1 1		japonicum structural	ľ			1
		muscle protein	ł	1		1
268	*****	paramyosin mRNA,	ł		1	
208	U11825	complete cds.	0.88	<none></none>	<none></none>	<none></none>
		Human DNA for	<u> </u>		(AL031907) hypothetical serine	1 1
269	D84278	CD38, exon 1			rich protein	1 1
207	D84278	Bovine lens aldose	0.68	3766363	[Schizosaccharomyces pombe]	3.0
1		reductase	1			_
270	M59755	pseudogene, 3' end.	0.67	MONTE		
	1.135735	pseudogene, J enu.	0.67	<none></none>	<none></none>	<none></none>
Į		Homo sapiens				
		skeletal muscle				
Į		voltage-dependent	İ			
		sodium channel alpha	i i			Í
J		subunit (SkM1)	ĺ		(Z86105) 1,4-beta-glucanase	
271		mRNA, complete cds.	0.65	2437819	[Anaerocellum thermophilum]	3.6
Ţ		Human type IV		2.5.012	(c.occitatii tilerinopiii)	J.U
j]:	sodium channel alpha	. 1		(Z86105) 1,4-beta-glucanase	ŀ
272	L01965	polypeptide	0.64	2437819	[Anaerocellum thermophilum]	3.5
		Danio rerio bone			the state of the s	
- 1		morphogenetic	1		(AE000720) formate	
		protein-4 (bmp4)	· 1		dehydrogenase alpha subunit	·I
273		nRNA, partial cds	0.63	2983532	[Aquifex aeolicus]	7.9
		Hylobates lar mucin			,	
22.		MUC1) gene, exons	1		(D79215) FGF-10 [Rattus	j
274	L41624 1	-6.	0.63	1517808	norvegicus]	0.91

	Nea	est Neighbor (BlastN vs	Genbank)	Negroet N	Jaiobhan (Dt. 1)	
SE	Q			.vearest iv	leighbor (BlastX vs. Non-Redundant	Proteins)
II.	ACCESS	ON DESCRIPTION	PVALUE	ACCESSIO		P VAL
\vdash					(U67956) coded for by C.	
275	AF03088	Fugu rubripes sushi retrotransposon gag polyprotein (gag) ar pol polyprotein (pol genes, complete cds Arabidopsis thaliana	nd)	1519696	elegans cDNA ykl26t9.5; code for by C. elegans cDNA ykl59h6.3; coded for by C. elegans cDNA ykl26f9.3; code for by C. elegans cDNA ykl59h6.5 [Caenorhabditis elegans]	d
276	U52909	U1 snRNP 70K protein gene, complete cds	0.62	. (NONE)	<none></none>	0.38
277	AF008192	Homo sapiens putative GR6 protein (GR6) mRNA, complete cds Human fatty acid	0.62	3800934	(AF100655) contains similarity to ser/thr protein kinases [Caenorhabditis elegans]	9.7
278	U17081	binding protein (FABP3) gene, complete cds	0.62	3617848	(AF049709) tyrosylprotein sulfotransferase-A; TPST-A	7.7
279	AB018340	Homo sapiens mRNA for KIAA0797 protein, partial cds	0.62	424044	VP5 protein - porcine rotavirus >gi 61355	7.7
280	Y00093	H.sapiens mRNA for leukocyte adhesion glycoprotein p150,95	0.62	1054945	(U38621) polyprotein [Tobacco	
81	M63138	Human cathepsin D (catD) gene, exons 7, 8, and 9.	0.62	136810	GLYCOPROTEIN M >gi 73791 pir WMBE51 UL10 protein - human herpesvirus 1 1- 473) [Human herpesvirus 11	4.5
32	X76056	N. sylvestris DNA for spacer region between 25S and 18S ribosomal RNA genes	0.62	2661176	>gi 221732 gnl PID d1002131 (U76671) putative cds Rhodobacter sphaeroides	2.0
3		B.taurus mRNA for ACTH receptor	0.62	4249552	(AB001075) galectin-2 related protein	2.0
1	S S	Rat F1-ATPase beta ubunit mRNA, 3'			transforming growth factor-beta type III receptor - chicken >gi 511843 (L01121) transforming growth factor-beta	
	-	nd.	0.62	2119692	type III receptor [Gallus gallus]	1.5

<u> </u>	Near	est Neighbor (BlastN vs	. Genbank)	Nearest Ne	ighbor (BlastX vs. Non-Redundant	Danie
SE	Q				Silvon-Redundant	Proteins)
ID	ACCESSI	ON DESCRIPTION	P VALUE	ACCESSION		P VALU
					ASPARTYLIASPARAGINYL	
					BETA-HYDROXYLASE	
1	j		ı	1	(ASPARTATE BETA-	1
	1	1	Ī	i	HYDROXYLASE) (ASP BET	A
	1		1		HYDROXYLASE) (PEPTIDE	-
	1	Homo sapiens	1	1	ASPARTATE BETA-	ľ
	1	SERCA3 gene, exor	ns	1	DIOXYGENASE) beta-	1
285	Y15724	1-7 (and joined CDS		2498164	dioxygenase (EC 1.14.11.16) -	ì
		Plasmodium		2490104	bovine >gi[162694 taurus]	0.52
ĺ	1	falciparum DNA ***	•		l	
	1	SEQUENCING IN	1		f	
	ł	PROGRESS ***	į.	1	İ	1
1	i	from contig 3-72,		1	HVPOTHETICAL PROTEIN	1
286	AL010142	complete sequence	0.62	3183206	HYPOTHETICAL PROTEIN KIAA0009 sapiens]	
				3103200	HYPOTHETICAL 63.5 KD	4e-07
	Ĭ		1	ł	PROTEIN ZK353.1 IN	i .
	ı		1	i	CHROMOSOME III	1
			1		>gi 1078903 pir \$44654	İ
		Mus musculus Stat3	1		ZK353.1 protein -	ĺ
		gene, 5'-flanking	1		Caenorhabditis elegans	l
287	AB008160	region and exon 1	1 1		>gi 289757 (L15313) putative	!
207	AB008160	partial sequence	0.62	466097	[Caenorhabditis elegans]	le-35
		1	1 1		(Z48585) similar to ATPases	10-35
- 1			ŀ		associated with various cellular	ſ
- 1			i 1		activities (AAA); cDNA EST	
1			i i		EMBL:Z14623 comes from this	
- 1			1		gene; cDNA EST	
- 1		Halomonas marina			EMBL:D75090 comes from this	1
- 1		gene for alginate			gene; cDNA EST	Į.
288	AB018795	lyase, complete cds	0.62	2055.00	EMBL:D72255 comes from this	1
T		Human DNA	0.02	3877493	gene; cDNA EST yk200e4	3e-46
- 1	•	sequence from	1	•		
- 1		cosmid E141E2, on			! · ·	ſ
		chromosome 22,			1	F
		complete sequence	- 1			1
289	Z69906	[Homo sapiens]	0.61	<none></none>	310375	
				VIVOITE >	<none></none>	<none></none>
ľ	i	Human clone ClITA-				
		8 MHC class [[ı			
<u>,, </u>		transactivator CIITA	1		(X79983) viral proteinase	
90	U18259	mRNA, complete cds.	0.61		(Pseudorabies virus)	
		S.tuberosum mRNA			(1 sessionables virus)	9.8
1		for inorganic		ľ		
91		phosphate ransporter, StPT1		lo lo	(U08884) protein VIII precursor	
V (0.61			

	_	.ta. No	earest	Neighbor (BlastN vs	. Genbank)	Nearest N	eighbor (BlaceV	
	SI	EQ			1 .	. TOLICSE 140	eighbor (BlastX vs. Non-Redundant P	roteins)
	1	D ACCES	1012	DESCRIPTION	P VALUI	ACCESSION	N DESCRIPTION	P VALUE
	-							IF VALUE
	29	92 U708	325	Rattus norvegicus 5 oxo-L-prolinase mRNA, complete co	ı	733543	(U23448) similar to genome polyprotein (SP:POLG_BVDVN, P19711);	
	29	3 L816	67	Homo sapiens (subclone 2_a9 from P1 H49) DNA sequence			(U80759) CAGH4 alternate open reading frame [Homo	4.4
	294	4 AE000	760	Aquifex aeolicus section 92 of 109 of the complete genome		2565087	sapiens] HOMEOBOX PROTEIN HOX- A3 (HOX-1.5) homeobox- containing transcription factor [Mus musculus]	3.3
	295	U5851		Mus musculus Rho- associated, coiled- coil forming protein kinase p160 ROCK-1 mRNA, complete cds			(L11275) selected as a weak suppressor of a mutant of the subunit AC40 of DNA dependant RNA polymerase I	2.6
- 1		1 3331		Human origin	0.61	295671	and III	1.5
	296	U27459	I I I	ecognition complex protein 2 homolog ORC2L mRNA, omplete cds isum sativum S-	0.61	200285	(M97900) putative open reading frame [Mus musculus]	0.66
	297	L36680	a s	denosylmethionine ynthase mRNA, 3'	0.61	2285790	(AB002086) p47 [Rattus norvegicus]	4- 12
	298	AE00067	se	quifex aeolicus ection 5 of 109 of e complete genome	0.61	3395782	(AF058446) histone macroH2A1.2 [Gallus gallus]	4e-12 6e-27
	299	AF086310	lei	omo sapiens full ngth insert cDNA one ZD51 F08	0.61	3646450	(AL031603) conserved hypothetical protein. [Schizosaccharomyces pombe]	
3	00	AJ009675	mI 3-r coe red	rotis ipsilon RNA for 3-hydroxy- nethylglutaryl enzyme A uctase	0.61	4176370	(AC005058) similar to calcium- independent phospholipase A2; similar to AC004392 (PID:g3367519) [Homo	8e-29
3(01	AC005577	chr cos cen con	mo sapiens omosome 19, mid F18382B, tromeric end, nplete sequence mo sapiens	0.60	<none></none>		2e-73
						CHOIRES	<none> <</none>	NONE>

-	Near	est Neighbor (BlastN vs	. Genbank)	Nearest N	leighbor (Bloss V	
SEQ					leighbor (BlastX vs. Non-Redundar	t Proteins)
ID	ACCESSI		P VALU	E ACCESSIO	DN DESCRIPTION	BVAL
_		Candida albicans				P VAL
1 .	ļ	topoisomerase type	1			
302	U40454	(CATOPI) gene,				1
1	040434	complete cds	0.60	<none></none>	<none></none>	<non< td=""></non<>
1 1		Emericella nidulans	1			·
J		mtDNA between		1	l	- 1
		h2/h5 and bh2/b2	j			1
i i		junctions, genes for	j	1		
1 1		ATPase subunit 6,		ł		ĺ
1		cytochrome oxidase	I		1	1
ľ		subunit 3, seven.		1	1	1.
		unidentified proteins	,	Ī		1
		twentyfour tRNA's		1		
303	J01390	and L-rRNA.	0.60	<none></none>	NONT	
		Plasmodium		4.0.112	<none></none>	<none< td=""></none<>
		falciparum RNA		İ	1	1
304	T 1 1 1 70	polymerase I gene,	1			1
307	L11172	complete cds. Caenorhabditis	0.60	<none></none>	<none></none>	ANONT
ı		elegans cosmid	1		1110112	<none:< td=""></none:<>
I		F39H11, complete				1
		sequence				į
		[Caenorhabditis	1 1			
305	Z81079	elegans)	0.00		1	j
$\neg \neg$		510 82.10	0.60	<none></none>	<none></none>	<none></none>
i		S.cerevisiae			MAJOR DNA-BINDING	
1	•	chromosome X			PROTEIN herpesvirus 1 (strain	ļ
		reading frame ORF			11) >gi 60327 (X64346) major	1
306	Z49627	YJR127c	0.60	118751	ssDNA-binding protein	
		Rattus norvegicus H-		110/51	[Saimiriine herpesvirus 2]	9.6
- 1		K-ATPase alpha 2				l .
		gene, alternatively	- 1	. •		1
07		spliced products and		•	(AF003086) PfSNF2L	
~		partial cds Methanococcus	0.60	2213862	[Plasmodium falciparum]	7.4
- 1		jannaschii section 18	Γ			/
- 1		of 150 of the	1		1	
08		complete genome	0.60		(D89240) unnamed protein	
		Methanococcus	0.60	1749688	product	5.7
j		annaschii section 55	1			
ſ		of 150 of the	i			
)9 t		omplete genome	0.60	2227421	(U97068) zonadhesin [Mus	l
			. 0.00	3327421	musculus]	4.3
		laemophilus ducreyi	1		(41.024401)	
.	li	poprotein gene,	1		(AL034491) conserved	
<u> </u>	J57817 c	omplete cds	0.60	4008577	hypothetical protein	j
				.000377	[Schizosaccharomyces pombe]	2.5

<u> </u>	Nea	rest Neighbor (BlastN	vs. Ge	nbank)	Nearest Mai	all (D)	
SEQ	:1				TACATEST TACT	ghbor (BlastX vs. Non-Redundant I	Proteins)
ID	ACCESS	ION DESCRIPTION	NC	P VALUE	1	DESCRIPTION	P VAL
	-				111	probable protein-tyrosine kinas	1. 171
311	X8070	D Paris C 17				(EC 2.7.1.112) RTK - Pacific	
	710070	H.sapiens G17 go Mus musculus (c	ene	0.60	422541	electric ray >gi 290858	٦.,
	i	P24) sde sees	ione			7 8.1220030	1.5
312	L42167	R24) rds genc, pa	irtial	_	İ	(AF033823) moira [Drosophila	1
	2,210,	cus		0.60	4220848	melanogaster)	1 00.
		Human hMSH6	I				0.51
313	U54777		.		1	(AF031087) mismatch repair	1
	30 1111	mRNA. complete Human mRNA fo	cas	0.60	2665637	protein MSH6 [Mus musculus]	5e-07
- 1		KIAA0232 gene,	·			i i i i i i i i i i i i i i i i i i i	36-07
314	D86985	complete cds				(U97006) No definition line	
	200703	Complete cas		0.60	1938462	found (Caenorhabditis aloona)	2e-07
- 1			- !	٠.		(USS376) coded for by C.	2e-07
1			- 1			elegans cDNA cm21e6; coded	
- 1			- 1			for by C. elegans cDNA	
- 1						cm01e2; similar to melibiose	
- 1		Rat liver mRNA fo	_			carrier protein	-
315	D43964	Kan-I, complete co				(thiomethylgalactoside permease	
		Rattus norvegicus	15	0.60	1280135	II)	5e-15
- 1		CTD-binding SR-li	1.0	ı			36-13
ł		protein rA4 mRNA				1	
16	U49058	partial cds	.	0.00		(U37500) RNA polymerase II	
$\neg \vdash$		U.ruddi		0.60	2145091	largest subunit [Mus musculus]	le-19
- 1		mitochondrial 12S	1				
17	X84388	ribosomal RNA	- 1	0.60	20	(Z70205) predicted using	
1		Caenorhabditis		0.60	3874247	Genefinder	2e-37
- 1		elegans cosmid	1	- 1			
18 /	AF125447	Y14H12B	1.	0.59			
		Hyoscyamus muticus	s ·	0.59	<none></none>	<none></none>	NONE>
- 1		clone cVS2		- 1			
-		vetispiradiene	1	- 1	1	1	
1		synthase mRNA,	1	- 1	ł	1	
9	U20189	partial cds.	1 0	0.59	NONT	1.	
		Human gastric H,K-	 	''''	<none></none>	<none> <</none>	NONE>
		ATPase catalytic	1	ı			
		subunit gene.	1	1	i i	1	
0 1		complete cds.	1 0	.59	NONE	1	
			 	·	<none></none>	<none></none>	NONE>
1		Helicobacter pylori					
1		(strain P1) comB and	1	- 1	- 1	. 1	ı
		omi/algA (partial)	1	ł	l l	ļ	
		genes, and partial				1	i
AJ		ORFI and ORF2	0.	59	NOVE		
			U.,	<i></i>	<none></none>	<none></none>	IONE>

	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Neis	nbor (BlastX vs. Non-Redundant F	
SE	Q				Total Vs. Non-Redundant P	roteins)
П	ACCESSIO		P VALUE	ACCESSION	DESCRIPTION	P VALU
<u> </u>		Mus musculus				IF VALU
1	j	transcription factor				+
		AP-2 (AP-2) gene,	1	1		
		alternative exon 1a,	1	Ĭ	i	1
1 22	,	and isoform 2, partia	l [i	(AC002332) hypothetical	1
32:	2 U17289	cds.	0.59	2459419	protein [Arabidopsis thaliana]	9.4
		S.cerevisiae				7.7
1		chromosome XIV	1	I	(Z67990) Similarity to Rat	
323	771466	reading frame ORF	1		amiloride-sensitive sodium	1
12	Z71466	YNL190w	0.59	3875542	channel beta-subunit	7.3
	i	Beet soil-borne virus				
1	1	genes for 13K, 22K	İ			· .
324	Z66493	and 48K proteins	0.50		cryV465 protein - Bacillus	
	200423	Homo sapiens	0.59	2119867	thuringiensis thuringiensis]	7.2
1	l l	prostasin mRNA,				
325	L41351	complete cds	0.59	720212	CRYSTALLIN J1C crystallin	
		S.lincolnensis gene	0.39	729212	[Tripedalia cystophora]	4.2
	1	for 16S ribosomal			(AFOSGS77) bill	
326	X79854	RNA	0.59	3702828	(AF056577) high mobility	
		Strongylocentrotus	- 5.52	3702028	group protein 1.2 HYPOTHETICAL PROTEIN	3.2
		purpuratus mRNA for			KIAA0129 product is novel.	
327	AJ223356	SuDp98 protein	0.59	2495704	[Homo sapiens]	2.5
220		H.sapiens mRNA for			(Y10027) transcription factor	2.3
328	X86019	PRPL-2 protein	0.59	1743341	TEF-1 [Mus musculus]	2.5
i		Xiphias gladius				
		creatine kinase gene,	l		(U69477) envelope glycoprotein	
329	U75528	partial cds	0.59	1045005	[Human immunodeficiency virus	
		partial cus	0.39	1845995	type 1] DNA POLYMERASE	2.4
		1 1	1		EPSILON SUBUNIT B DNA-	
		1	1		directed DNA polymerase (EC	1
		j			2.7.7.7) II chain B - yeast.	
				•	(Saccharomyces cerevisiae)	
	à		. [>gi 786319 (U25842) DNA	ſ
			1		Polymerase epsilon, subunit B	j
		Homo sapiens	i	i	(Swiss Prot. accession number	ļ
		chromosome 5, PAC	ľ		P24482) [Saccharomyces	i
330		clone 202e13	0.59	2506366	cerevisiae	1.4
Į		Rat receptor-linked				17
221		protein tyrosine	l		(X96713) collagen [Globodera	
331	L19180	phosphatase	0.59		pallida]	1.1
ļ	l,	Listeria	1			
ĺ		monocytogenes secA	i		1	
332		gene, complete cds.	0.50		AF016415) No definition line	4
		one, complete cus.	0.59	2291129	ound [Caenorhabditis elegans]	0.83

<u> </u>	Near	est Neighbor (BlastN vs	. Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant F	rotains)
S	EQ				T Ten redundant i	Toterns)
L	D ACCESSI		P VALUE	ACCESSION	DESCRIPTION	DVALLE
<u> </u>		Xenopus laevis			(AL031124) hypothetical	PVALUE
		syndecan-2 mRNA,			protein SC1C2.25c	
3	33 U24433	complete cds.	0.59	3355692	[Streptomyces coelicolor]	0.64
İ					(acceptomyces coefficient)	0.64
		Drosophila	i	ł		İ
	1	muscarinic				1
33		acetylcholine recept	or		(M76546) hydroxyproline-rich	1
133	4 M23412	mRNA, complete co	s. 0.59	168237	protein [Helianthus annuis]	0.22
1					protein [Helianthus annuus] HYPOTHETICAL 67.5 KD	0.22
1	1				PROTEIN IN PRPS4-STE20	
	1				INTERGENIC REGION	l i
	j			• ,	>gi 626567 pir S46825	
	ł	Symanhau			hypothetical protein YHL010c -	1 1
1	ł	Synaphea media	.]		yeast (Saccharomyces	1 1
1	Ï	chloroplast atpB-rbcl	-		cerevisiae) >gi 2289881	i i
			1		(U11582) No definition line	i i
335	AF060729	region, partial			found [Saccharomyces	
1 233	AL 000729	sequence	0.59	731596	cerevisiae]	0.16
ĺ	1	1	1 1			
1	İ	Xanthobacter	1 1			ı
1	1	autotrophicus	1 1			
	ł	transcriptional	1		1	
1	I	activator AldR (aldR)	1 1		1	ŀ
	1	gene, partial cds; and	1 1		1	
	1	NAD-dependent	i i		PERIAXIN	í
		chloroacetaldehyde	1 1			- 1
		dehydrogenase (aldB)			>gi 2143901 pir 158157 periaxin	
336	AF029734	gene, complete cds	0.59	2498801	- rat >gi 505297 (Z29649)	
				2470001	periaxin [Rattus norvegicus] HYPOTHETICAL 34.3 KD	0.13
	,				PROTEIN IN TAF145-YOR1	1
	1				INTERGENIC REGION	ı i
	ł		1		>gi 2131717 pir S64612	
	ł	1			hypothetical protein YGR277c -	
	l	1	1		yeast (Saccharomyces	
		!	İ		cerevisiae)	j
			i		>gi 1323505 gnl PID e243248	
227	3/0 400-	C.reinhardtii LI818r-	Į.		(Z73062) ORF YGR277c	į.
337	X95307	1 genc	0.59	1723781	[Saccharomyces cerevisiae]	le-04
		Dictyostelium	7		HYPOTHETICAL 43.3 KD	10-04
ı		discoideum tRNA-	1		GTP-BINDING PROTEIN IN	
338		Glu-GAA gene, clone	1		DACB-RPMA INTERGENIC	1
220	M24572	yGluGAA8.	0.59	1176186	REGION >gi[606121 coli]	3e-06
I		Human h) (CI)	1			
339	• · • • • • • • • • • • • • • • • • • •	Human hMSH6 gene.	0.50	J.	(AF031087) mismatch repair	Ì
	3,3,35	CAUL 2	0.59	2665637	protein MSH6 [Mus musculus]	5e-07

\vdash	Neare	st Neighbor (BlastN vs.)	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant	
SE	Q				Endor (Blasta Vs. Non-Redundant	Proteins)
IL	ACCESSIC		P VALUE	ACCESSION	DESCRIPTION	P VALU
<u> </u>		Escherichia coli				1. VALU
340	D007	genomic DNA. (25.2	1			
341	D90747	25.6 min)	0.59	134286	DOLICHOL KINASE	6e-08
ļ			.]		major centromere protein.	De-08
341	105711	Human desmoplakin	1		CENP-B [human, Peptide, 594	1
- 	J05211	mRNA, 3' end.	0.59	246796	aa]	4e-08
1		I olive and the	ļ		KINESIN LIGHT CHAIN	10-08
ľ	1	Loligo pealii kinesin	l	Ĭ	(KLC) sea urchin	
342	L24441	light chain mRNA,		ĺ	(Strongylocentrotus purpuratus)	İ
3 12	1.24441	complete cds. Human cardiac alpha-	0.59	547800	>gi 161530	5e-14
ı	į		ł			1 30 17
	I	myosin heavy chain		'		1.
343	M25140	(MYH6) gene, exons 2. 3 and 4.				1
3 +3	14123140	Homo sapiens	0.58	<none></none>	<none></none>	<none></none>
		(subclone 9_h2 from				
	1	P1 H21) DNA		l	1	
344	L81932	sequence			1	1
	201752	Homo sapiens full	0.58	<none></none>	<none></none>	<none></none>
	1	length insert cDNA	j			
345	AF087966	clone YU51G04	0.50			
		H.sapiens flow-sorted	0.58	<none></none>	<none></none>	<none></none>
	i	chromosome 6 TaqI	ł			
		fragment,	1		•	
346	Z 78574	SC6pA10G11	0.58	ALONIE	1	
		Blattella germanica	- 0.30	<none></none>	<none></none>	<none></none>
		allatostatin	ľ			
		neuropeptide	- 1		1	
		precursor, gene,	i			
347		complete cds	0.58	<none></none>	<none></none>	
		Homo sapiens Cdc7		4.01122	SNOINES	<none></none>
		(CDC7) mRNA,				1
348	AF015592	complete cds	0.58	<none></none>	<none></none>	NO. T
	i,	Markana .			CROIVES	<none></none>
- 1		Methanosarcina	i			Í
	ſ°	barkeri atp operon:	1			- 1
- 1		ATP synthase beta	ł			ļ
- 1		subunit (atpD), ATP	- 1		1	f
- 1		ynthase epsilon	j			- 1
ı		ubunit (atpC), ATP	i			- 1
		ynthase gene l	1	i		
19		atpI). ATP synthase	. 1	ļ	(AC004136) putative DNA	
''	AF028006 a	subunit subunit (0.58	3184291	polymerase III gamma subunit	9.4
- 1	١.	S				
		lus musculus gene	I	1		
ı		or pancreatic trypsin, omplete cds	- 1	1,	(AF056704) synapsin IIIa	1
0	ARDI7022		0.58	3170561		

	Neares	t Neighbor (BlastN vs.	Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant Pr	
SE			T	THE STATE OF THE S	BlastA vs. Non-Redundant Pi	oteins)
II	- 1		P VALUE	ACCESSION	DESCRIPTION	PVALUE
-		Dictyostelium				I. VALUE
351	AF081585	discoideum developmental protein DG1110 (DG1110) gene, partial cds	0.58	105417	basic proline-rich peptide IB-8a	9.2
352	AFORCOO	Homo sapiens full length insert cDNA			hypothetical protein - African swine fever virus (strain Malawi Lil-20/1) >gi 450758 (X71982) myeloid differentiation antigen homologue [African swine fever virus] >gi 903686 (M95672)	
332	AF086322	clone ZD53E01	0.58	93026	unknown protein	7.1
353	AF088025	Homo sapiens full length insert cDNA clone ZC19C04 Human mRNA for	0.58	2384644	(U92805) thrombospondin-3 [Xenopus laevis]	7.0
354	AB002339	KIAA0341 gene, partial cds Methanococcus	0.58	2135587	M130 antigen (cytosolic variant 2) - human	5.4
355	U67548	jannaschii section 90 of 150 of the complete genome	0.58	2911094	(AL021957) hypothetical protein Rv2174	4.2
356		Homo sapiens receptor tyrosine kinase (ERBB4) gene, complete cds.	0.58	461922	PYRUVATE DECARBOXYLASE (8-10 NM CYTOPLASMIC FILAMENT- ASSOCIATED PROTEIN) (P59NC) 4.1.1.1) - Neurospora crassa >gi 293948 (L09125) pyruvate decarboxylase [Neurospora crassa] >gi 1655909.	4.2
357		Bacillus subtilis sigma 43 operon with P23-dnaE-rpoD genes (dnaE for DNA primase, rpoD for RNA polymerase)	0.58	1323704	(U55387) similar to C. elegans F38E1.9 gene product encoded by GenBank Accession Number U41996 [Cricetulus griseus]	
358	I c r r	Desulfovibrio /ulgaris rbo gene for lesulfoferrodoxin and ub gene for ubredoxin, complete ds			(AC004680) putative protein	4.1
	2,0419	из	0.58	3420047	kinase [Arabidopsis thaliana]	2.4

	Neares	t Neighbor (BlastN vs.)	Genbank)	Nearest Neis	≥hbor (BlastX vs. Non-Redundant P	
SEC	2				- Verasor vs. From Redundant P	roteins)
ID	ACCESSIO		P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Human DNA				I. VALUE
1	,	sequence from				
1	ı	cosmid B20F6 on	1	Ì	1	1
ļ	i	chromosome 22,	1	Į.	1	
	1 .	complete sequence		Į.		1
359	Z82174	[Homo sapiens]	0.58	2145455	(Y07866) catalase-peroxidase	2.4
	1	F.solani STI35				
360	16225.5	protein gene.			(AL021897) hypothetical	ł
360	M33642	complete cds. Mus musculus	0.58	2896706	protein Rv1069c	2.4
1	1		1			
1	į –	transforming growth	ļ	i	(Z81038) predicted using	1
		factor alpha (TGF		٠.	Genefinder; cDNA EST	· ·
361	U64873	alpha) gene, partial			yk488a2.5 comes from this gene	ļ
101	004873	cds Macrophthalmus	0.58	3874437	[Caenorhabditis elegans]	1.8
1		banzai mitochondrial				
1		DNA for 12S and	l		1	
		16S rRNA, partial	ľ			
		and complete			(4 (224626))	
362	AB002132	sequence	0.58	2960022	(AJ224676) rho type GEF	
		Caenorhabditis	0.50	2500022	[Drosophila melanogaster]	1.8
		elegans MutS				
li		homolog (msh-5)		_	(U75869) Omp22 [Helicobacter	
363	AF070070	mRNA, partial cds	0.58	4098205	pylori]	1.8
		Staphylococcus				
		epidermidis plasmid	i			ł
		pIP1629 mobilization	1		1	i
364	AF045240	protein (mobC1),				- 1
304		(orf69-1), (mobA1), H.sapiens Wilms	0.58	4218117	(AL035353) protein (fragment)	0.62
l j		tumor gene 1, exons 8	I			
365		and 9	0.58	2221050	(U88211) unknown [Gallus	
			0.58	2331059	gallus]	0.62
· 1		. 1			1	l
	į.	Moraxella catarrhalis	1		1	ı
i		strain 4223 transferrin	1		1]
1		oinding protein A	ł			
_ ,		(tbpA) and transferrin	- 1		FIBER PROTEIN	1
`		oinding protein B	1		>gi 74229 pir ERADFM fiber	1
		tbpB) genes,	1	•	protein - mouse adenovirus I	
366		complete cds; and			>gi 209758 (M30594) fiber	
300		inknown gene	0.58	120155	protein [Mastadenovirus musl]	0.27
- 1		Iuman mRNA for	ı			
367		CIAA0273 gene,	2.50	224	(U94177) androgen receptor	ŀ
 +	207403	omplete cds	0.58	3861477	[Pan troglodytes]	0.12
1	N	lus musculus ninein	1	1	(15110003) 37 1 5 1 1	
368		iRNA. complete cds.	0.58		(AF118223) No definition line	
		topiete eds.[0.50	411000	found [Arabidopsis thaliana]	0.004

200

	Neares	st Neighbor (BlastN vs.	Genbank)	Nearest Nair	ghbor (BlastX vs. Non-Redundant F	
SE	Q			TOWN CONTINUES	thou (Blasex vs. Non-Redundant F	roteins)
	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU
-		CD68=110kda				1
	1	transmembrane	1			1
1	ſ	glycoprotein [humar				
	1	promonocyte cell lin		·		
		U937, mRNA, 1722	٠	1		
36	9 S57235	nt]	0.50	207250	(U96113) WWP1 [Homo	
	937,233		0.58	2072501	sapiens]	le-04
	ſ	Mus musculus	İ	ľ	(Decough) =	
1		serotonin1A receptor	. [I	(D63481) The KIAA0147 gene	}
370	U39391	mRNA, complete cd		1460076	product is related to adenylyl	1
			0.38	1469876 ·	cyclase. [Homo sapiens]	le-07.
		Monkey B-				
1	1	lymphotropic	• •		1	ĺ
1	ı	papovavirus genes fo	r			1
	1	VP-1, 2, 3 and large	1			ł
1	1	T antigen, complete	1			
	ı	and partial cds, strain	1		1	
1	i	LPV-76 > ::				
1	İ	gb M14494 PPMVP1				
	1	M Monkey B-	1 1			
		lymphotropic	i .		ĺ	
l	1	papovavirus mutant			1	
	j	(LPV-76) PstI B	í i			
l	1	fragment encoding	1			
, , , ,		VP1, VP2, VP3 and	1		(AJ001774) vanadium	İ
371	D00056	T-antigen.	0.58	2462069	chloroperoxidase	le-08
		A			HYPOTHETICAL 43.8 KD	10-00
		Amsacta	- 1		PROTEIN IN NCE3-HHT2	
		entomopoxvirus	ł		INTERGENIC REGION	
	1 i	spheroidin gene,	ŀ		>gi 2131871 pir S62957	
		complete cds, and			hypothetical protein YNL035c -	
	í ' I	four vaccinia related	1		yeast (Saccharomyces	
		orfs. > ::	J	•	cerevisiae)	İ
		gb I16670 I16670	J		>gi 1301880 gn1 PID e239670	Į.
372		Sequence 1 from	2.50		(Z71311) ORF YNL035c	- 1
7,2		patent US 5476781 gloo-S=growth-	0.58	1730722	[Saccharomyces cerevisiae]	8e-14
		associated protein				
373		GAP-43 homolog	0.50	0.000	(AF037168) DnaJ homologue	. [
	2,23,7	Solomon Ce- TVC	0.58	2689720	[Arabidopsis thaliana]	7e-14
	l ₂	Tetraodon fluviatilis	1		HYPOTHETICAL 34.9 KD	
ſ		imyloid precursor	f		PROTEIN C57A10.11C IN	
		protein mRNA.	. 1		CHROMOSOME I	ľ
374		complete cds	0.58		>gi 2058378 gn1 PID e314002	į
			0.50	3219938	pombe]	5e-22

	Neares	st Neighbor (BlastN vs.	Genbank)	N		
SE	Q.	Theighbor (Blastia Vs.	Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundar	nt Proteins)
n		N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-		F9 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				I VALUE
		Filobasidiella				
	1	neoformans		1		- 1
	1	translation elongation	n	ľ	1	
		factor EF1-alpha		}	ì	1
37	5 U81803	(CnTEFI) mRNA, complete cds	1 050		.]	i .
	081803	Candida albicans	0.57	<none></none>	<none></none>	<none></none>
		ATCC 18804, CBS	1	<u>[</u>		
	1	562 peptide	1		İ	
		transporter gene,	ĺ		·	1
376	U09781	complete cds.	0.57			
		Homo sapiens	0.57	<none></none>	<none></none>	<none></none>
1	1	(subclone 4_b10 from	,			
		BAC H102) DNA]		1	1 1
377	AC002143	sequence	0.57	<none></none>	2107	
		Tetrahymena		4101122	<none></none>	<none></none>
	i	thermophila RR			1	1 1
1		internal deletion] [
378	U23442	sequence.	0.57	<none></none>	<none></none>	<none></none>
	1	Mus musculus	. 1			CHOINES
	ł	transcription factor				1 1
	ł	AP-2 (AP-2) gene,		•	1	ļ
	1	alternative exon la.	•		<u> </u>	1
379	U17289	and isoform 2, partial				
	017289	cds.	0.57	<none></none>	<none></none>	<none></none>
		Buzura suppressaria	j			
	i i	nuclear polyhedrosis	i			
		virus gene for				1 1
380		polyhedrin protein	0.57	<none></none>		
			0.57	CHOILES	<none></none>	<none></none>
		Homo sapiens 5T4	j	Ţ		
		oncofetal trophoblast	. 1	j		1 .1
381	AJ012159	glycoprotein gene	0.57	<none></none>	<none></none>	NONE
		H.sapiens simple			CHONES	<none></none>
382		DNA sequence region	ĺ			
202	X76571 c	clone wg la8.	0.57	<none></none>	<none></none>	<none></none>

	Nec	rest Neighbor (BlastN	vs. Genbank)	Nearest N	eighbor (Planty	
S	EQ			10203111	eighbor (BlastX vs. Non-Redundani	Proteins)
	D ACCESS	ION DESCRIPTIO	N P VALUE	ACCESSIO	N DESCRIPTION	
				1 110023310	N DESCRIPTION	P VALU
- 1		Vibrio cholerae				
- 1	I	pathogenicity isla	nd, ∫	1		
	i	putative transposa	ıse,	1	1	
		aldehyde		1	1	ł
	ı	dehydrogenase	ļ	j		ł
- 1	- 1	(aldA), toxR-	į	ĺ	1	
ł	ı	activated gene A	ł			
	j	protein (tagA), putative inner	1	1		
- 1	· ·	membrane protein,	1	1	J	1
	1	and putative zinc	· }			j
	ı	metalloprotease	İ	1		1 .
	1	genes, complete cd		1	1	1 .
38	3 AF03443	4 and	· 1			
		Mus musculus gene	0.57	<none></none>	<none></none>	<none></none>
	ı	for TESP4, comple				
384	4 AB01703	1 cds	0.57	<none></none>		
		S.hispidus	0.57	KNOMES	<none></none>	<none></none>
1	i	mitochondrial DNA				
1_	.	for SSU ribosomal				1 1
385	X89788	RNA gene	0.57	<none></none>	-NONE	1
	1	Rat progesteron		4.01.62	<none></none>	<none></none>
386		receptor gene, 5'	1 1		(AE001251) femA protein.	1 1
300	L16921	untranslated region.	0.57	3323116	putative [Treponema pallidum]	8.9
1	ı	Hama and	1		(S48157) DNA polymerase-	8.9
l		Homo sapiens interferon regulatory	1 1		primase 180 kda subunit	1
387	AF027292	factor 6			[Drosophila melanogaster,	1 1
	12 02/232	Cicer arietinum	0.57	259790	Peptide, 1490 aa]	6.7
	1	mRNA for	1		hypothetical protein YDR-146w -	
388	AJ012581	cytochrome P450	0.57	2121400	yeast CAI: 0.11 [Saccharomyces	1
		7.50	0.57	2131498	cerevisiae]	5.3
	i	Human transfer RNA				
	j	Met (TRMEPI)	1 1		1	
200	J	pseudogene, complet	e		(AF070935) GABA receptor	1
389	L15363	gene	0.57	3228680	subunit [Musca domestica]	
	,	Helicobacter pylori			(U97008) weak similarity to	5.2
		26695 section 3 of	1 1		family 1 of G-protein coupled	1
390	4 F000636	134 of the complete	1		receptors [Caenorhabditis	l l
390	AE000525	genome	0.57	1938478	elegans]	4.0
		Amblyomma	Ι Γ			
		americanum	j <i>1</i>		1	ı
1		ecdysteroid receptor	1 1	•	1	ſ
		(AamEcR) mRNA.	1		1	ļ
391	AF020189	3'UTR. region 1	0.57	2072224	(U94875) p40 [Borna disease	ł
			9.57	2072224	virus]	4.0

-	Nea	rest Neighbor (BlastN vs	. Genbank)	Nearest N	leighbor (Pl W	
SEC	ર			T. Garest IV	leighbor (BlastX vs. Non-Redundant l	Proteins)
ID	ACCESS	ION DESCRIPTION	P VALU	- 1		
		Human UbA52 gene	1 1 1 1	E ACCESSIO	DESCRIPTION	P VAL
		coding for ubiquitin				
1	I	52 amino acid fusion	.			
392	X56991		0.57	3000110	(AL022121) hypothetical	1
1			1 0.57	2960113	protein Rv3689	4.0
1	1	Plasmodium	1	1	CYTOCHROME P450 2C14	
I	1	falciparum DNA ***	•		(CYPIIC14) phenobarbital-	
i	1	SEQUENCING IN		i	inducible, hepatic - rabbit P-450	Y .
	j	PROGRESS ***	1	ı	[Oryctolagus cuniculus]	i
200		from contig 4-81,	l l	į.	>gi 358265 prf 1306317A	i
393	AL01026	O complete sequence	0.57	117233	cytochrome P450 [Oryctolagus	l
	1			117233	cuniculus]	3.9
	I	Xenopus laevis	1	1	GASTRULA ZINC FINGER PROTEIN XLCGF44.2	
	l	gamma-crystallin		1	Soil857361=:-IISO6571 G	,
394	Maggar	(gcry3) gene,	ŀ		>gi 85736 pir S06571 finger protein (clone XIcGF44-2) -	
-354	M99581	complete cds.	0.57	141647	African claused for 45	
ł		Drosophila			African clawed frog (fragment)	3.0
- 1		melanogaster seven in	1	į.		
395	M38384	absentia mRNA,			(U80454) T16A1.1	
	14130304	complete cds.	0.57	1707127	[Caenorhabditis elegans]	2.0
ı		Haemophilus	ı		e e e e e e e e e e e e e e e e e e e	3.0
1		influenzae Rd section		1	IRON(III)-TRANSPORT	
- 1		110 of 163 of the			SYSTEM PERMEASE	
396	U32795	complete genome	0		PROTEIN SFUB >gi 152861	
		Klebsiella	0.57	1173433	(M33815) protein (sufB)	.2.3
		pneumoniae nifX,			, , , , , , , , , , , , , , , , , , , ,	٠.٠
		nifU, nifS, nifV and	ı			
397	X12600	nifW genes	0.57	22224	(AL021925) hypothetical	
			0.57	2909562	protein Rv2256c	1.4
		Homo sapiens mRNA	ı		insect-stage-specific protein -	
.		for KIAA0626	1		Trypanosoma cruzi >gi 162099	
98	AB014526	protein, complete cds	0.57	493300	(M65021) insect stage-specific	1
		Rhodococcus fascians	<u> </u>	482390	antigen	0.61
		strain NRRL-B-	1			
ł	Í	15096 hypothetical	1			
<u>, </u>		protein gene,	Í		(AFO2 ISOS	ı
9 4	AF063587	complete cds	0.57	4104321	(AF034582) vesicle associated	
	ĺ			7104321	protein [Rattus norvegicus]	0.46
0		Guinea pig estrone			alphy/hata at ' t'	
<u> </u>	L11117 S	sulfotransferase gene.	0.57	82584	alpha/beta-gliadin precursor	- 1
				02007	(clone A212) - wheat	0.35

<u> </u>	Near	est Neighbor (BlastN vs.	Genbank)	Nearest Nei	ohbor (BlackV V	
SEQ				TO LOSE THE	ghbor (BlastX vs. Non-Redundant	Proteins)
ID	ACCESSI	ON DESCRIPTION	P VALUE	ACCEPTAGE		
			TIVALUE	ACCESSION	DESCRIPTION	P VALL
		Mouse complete gene	el			
		for a mouse kallikrein	.]			
- 1		gene. Genes are mGK		į.		
ĺ		l (complete gene)	1	ŀ		
ď	l	and mGK-2 of	İ	1	1	
ł		hormones, e.g.,	ĺ	1	-}	1
		grow > ::			NE VOT TAR THE T	
- 1		gb J00390 MUSKAL	1	Í	NUCLEAR HORMONE	1 .
I		07 Mouse pseudo-	ĺ	f	RECEPTOR NOR-2 receptor	1
		kallikrein 2, exons 4	ĺ	ĺ	[Rattus norvegicus]	1
_ 1		and 5, and kallikrein			>gi 1583604 prf 2121281A	1
401	V00829	I gene, complete cds.	0.57	2500916	NOR-2 protein [Rattus	
- 1				2300310	norvegicus]	0.20
f		Chicken mRNA for			(11/0053)	
		beta-2 subunit of			(U40953) similar to matrin F/G	İ
	•	neuronal nicotinic	- 1		(SP:Q00910) containing C4- type zinc-fingers	ĺ
102	X53092	acetylcholine receptor	0.57	1072256	[Caenorhabditis elegans]	
- 1					(Z81035) predicted using	0.031
- 1			1		Genefinder; Similarity to	
			I		dehydrogenases; cDNA EST	
			i		EMBL:D65800 comes from this	
- 1		1	ı		gene; cDNA EST	
		1	- 1		EMBL:D76184 comes from this	
- 1		Ovis ovis granulocyte			gene; cDNA EST	i
- 1		colony stimulating	- 1		EMBL:D69322 comes from this	
03	L07939	factor			gene; cDNA EST	
7		Colletotrichum	0.57	3874345	EMBL:C08158 comes f	-3e-07
- 1		gloeosporioides	ŀ			
ŀ		CAP20 (cap20) gene,				
4	U18061	complete cds.	0.57	2014605	(AC003974) putative ubiquitin	ľ
				2914695	specific protease	9e-08
1			1	·	TUMOR NECROSIS FACTOR,	
			I		ALPHA-INDUCED PROTEIN	· 1
	1	}	1.	j,	3 (PUTATIVE DNA BINDING	ł
-			1	l;	PROTEIN A20) (ZINC	- 1
1]:	L.japonicus mRNA	- 1	,	FINGER PROTEIN A20)	
1	Į:	for small GTP-	1		>gi 107549 pir A35797	1
		binding protein,	j	.	probable DNA-binding protein A20 - human >gi 177866	- 1
	273955 II	RABLIG		12	540 - πuman >gil177866	T I

	Near	est Neighbor (BlastN vs.	Genbank)	Nearest Neigh	hor (Blast V us No. 2)	
SE			7		bor (BlastX vs. Non-Redundant F	roteins)
ID	~	ON DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-	 					T
1	ı	1.	1	1	(Z77660) Similarity to Human	——
	l	1	Į.	i	enoyl-CoA hydratase	
1	1	}			(SW:ECHM_HUMAN); cDNA	1
	j	i	İ		EST EMBL:T00611 comes	
	1				from this gene; cDNA EST yk203d10.3 comes from this	1 1
	1	Petunia grp-1 gene			gene; cDNA EST yk203d10.5	1 1
	1	for glycine-rich	1		comes from this gene; cDNA	1 1
406	X04335	protein	0.57	3876901	EST yk457h5.3 comes from t	
	1	Rattus norvegicus S-			251 yk457115.5 comes from t	le-27
1	i	adenosylmethionine	1 1	٠,	ĺ	
	ľ	decarboxylase			1	1 1
407	FIADZIA	(AMDP2)	1 1		1	i i
407	U40718	pseudogene S.cerevisiae SSD1	0.56	<none></none>	<none></none>	<none></none>
		protein gene,	1			
	j	complete cds. > ::	1			
		gb AR013983 AR013	1 1			
		983 Sequence 8 from	1 1			
408	M60318	patent US 5773245	0.56	<none></none>	-NONTE	
		Nicotiana tabacum		T.OIVE2	<none></none>	<none></none>
		blp4 mRNA for	1 1		 	
400	7550	luminal binding	İ			
409	X60057	protein (BiP)	0.56	<none></none>	<none></none>	<none></none>
		Homo sapiens full			,	
410	AF085930	length insert cDNA		ŧ		1
***	717 003930	clone YR55A09 Plasmodium	0.56	<none></none>	<none></none>	<none></none>
		falciparum DNA ***	£ .			
		SEQUENCING IN	1	•	1	j
ı	•	PROGRESS ***	- 1		. 1	ł
		from contig 3-102,				
411	AL010189	complete sequence	0.56	<none></none>	<none></none>	<none></none>
		Murine G-CSF gene				CIVOINES
ł		for granulocyte	j	ĺ		i
412	X05402	colony stimulating	1			
***	703402	factor precursor Rattus norvegicus	0.56	<none></none>	<none></none>	<none></none>
ł		regulator of G-protein	I			
- 1		signalling 12	1	1		j
		(RGS12) mRNA,	1	I		Į.
113		complete cds	0.56	<none></none>	-NOVE	
		Human		ATOMES .	<none> <</none>	NONE>
		papillomavirus strain	• 1	j	Į.	
	****	RTRX7 complete	1	1		
14	U85660	genome	0.56	<none></none>	<none></none>	NONE>
						4.0

<u> </u>		t Neighbor (BlastN vs. (Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant F	rotair -
SEQ					10.12.24 VS. (VOII-Redundant F	Toteins)
ID	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	
			T T		DESCRIPTION .	P VALU
	Í	M. javanica				
	i	mitochondrion	ļ	1	•	
		ATPase 6, and		1		
415	X57626	putative tRNA-f-Met		1		1
	A37026	and tRNA-His genes Sus scrofa \$100C	0.56	<none></none>	<none></none>	<none></none>
416	AB003363	gene, complete cds	0.55			1
	112003303	gene. complete cas	0.56	<none></none>	<none></none>	<none></none>
		Danio rerio DANA		l	(AE001062) conserved	
417	L42291	element, intron 4.	0.56	3650000	hypothetical protein	
		1	0.30	2650002	[Archaeoglobus fulgidus] L-LACTATE	8.7
		·	l		DEHYDROGENASE	
			٠.	•	(IMMUNOGENIC PROTEIN	!
- 1					P36) >gi 479296 pir \$33362 L	
ı		Mus musculus			lactate dehydrogenase (EC	
		leukocystatin gene,			1.1.1.27) - Mycoplasma	
418	AF031826	complete cds	0.56	462493	hyopneumoniae	<i>(</i> 7
		Pennisetum glaucum			пуорнеатопас	6.7
419	TH TOCO	Ac-like element,	ſ		ESCARGOT/SNAIL PROTEIN	
719	U17068	AcL2.	0.56	399449	HOMOLOG	6.7
1		H.sapiens mRNA encoding GPI-	l		HYPOTHETICAL 8.7 KD	0.7
- 1		anchored protein	l		PROTEIN (READING FRAME	
120	Z48042	p137	0.56		D) >gi 76316 pir QQSA7C	
		Choristoneura	0.36	141232	hypothetical protein E-74	6.7
- 1		fumiferana	ľ		ACETYLCHOLINE	
ĺ	1	entomopoxvirus	j		REGULATOR UNC-18	
ł	Į.	nucleotide	J		>gi 480359 pir S36747	i
- 1		triphosphate	+		acetylcholine regulator unc-18 -	1
	ļ:	phosphohydrolase I	i		Caenorhabditis elegans	ľ
l	1 222	NPHI) gene,	1		>gi 247392 bbs 100294 putative	
21	AF027657	complete cds	0.56	464999	acetylcholine regulator unc-18	5.1
- 1		·			URACIL-DNA	
- 1	1,	January D.V.	1		GLYCOSYLASE (UDG)	i
		Homo sapiens mRNA			herpesvirus 2 >gi 695219	ł
22		or MEGF7, partial			(U20824) uracil DNA	
1		ua	0.56	1718033	glycosylase	5.1
- 1	ł	1	l		COMMON PLANT	
- 1	i		1		REGULATORY FACTOR	j
- 1	lx	maculatus NGF	·		CPRF-1 >gi 515621 (X58575)	- 1
1		ene for nerve growth			light-inducible protein CPRF-1	
3		ctor	0.56	1169081	[Petroselinum crispum]	
			0.30	1109081	>gi 1498301 (U46217) CPRF1	3.8

<u> </u>		IVE	arest	Veighbor (BlastN	vs. G	enbank)	Neare	et Mai	abb - (D)		
	SEQ			}				reare	21 1ACI	ghbor (BlastX vs. Non-Redunc	iant Pr	roteins)
-	ID A	CCES	SION	DESCRIPTIO	N	PVA	LUE	ACCES				
				Rat transcriptiona						DESCRIPTION		P VAL
	ı			repressor of myeli	۱ ۱							
	- 1		- 1	specific as (Sc	n-							
4	24	M727	l.	specific genes (SC	ויש		- 1			(U01849) ORF2 [Trypanos		
			·	mRNA, complete	cds.	0.5	5	50102	27	bruceil	oma	
1	- 1		ı				- 1			CHORISMATE MUTASE		2.3
1	- 1				- 1		- 1			(CM) / PREPHENATE	- 1	
-	- 1		l c	aenorhabditis		•	- 1			DEHYDRATASE (PDT) (F	2.	
1	- 1		- 1	legans cosmid			- 1			PROTEIN)	1	
1	ı			67D11A, comple	e		ı			>gi 281791 pir S26053		
1	- 1			quence	٦		- 1			chorismate mutase (EC 5.4.0	99.51	
1	_		[[Caenorhabditis						P / prephenate dehydratase /	FC	
42.	3 AI	L0238		egans]	-	0.56	.	2667-		4.2.1.51) - Erwinia herbicola		
1	1		T		+	0.20	+	266771		>gi 43344		2.3
1			- 1.				- 1			ATD		
	- 1		Sc	histosoma mansoi	ıi 📗					ATP synthase chain 6 -	ı	
l	1		gу	necophoral canal						Platymonas subcordiformis	Í	
426		47862		otein mRNA,	1		- 1			mitochondrion >gi 633582		
720	1-0	4/862	COI	mplete cds	\bot	0.56		2147138		(Z47797) ATP synthase subu	nit	
	1		الت.							6 [Platymonas subcordiformis	<u>;] </u>	2.3
	1		200	man germ line	1		1					
	1		bla	e homologous to	1		i				- 1	
	1		Onc	ogene T24 (Gene	1		1		ſ		- 1	
	ł		cod	e c-Ha-ras-1) with	1		1		- 1	(U60289) receptor protein	- 1	
427	L Vo	0574	four	exons.		n \- 4	1		·],	tyrosine phosphatase psi [Hom		- 1
				evis H1(0)-1	-	0.56	-	1518672		sapiens]	"	1.7
428	Z7.	1502	gene		ı	0.56	1			D90899) ferrichrome-iron	+	1./
			T-		-	0.36	+-	1651674	r	eceptor		1.3
ا ۵۵			R.no	rvegicus renin			1		K	AF045141) putative serine	+	
429	M37	7278	gene	exons 1-9.	(0.56	1	2057010	P	roteinase [Scirpophaga		1
					`		 	2853019	<u>[ir</u>	ncertulas]		1.0
- 1			Then				l			•		
			therm	ophilus polA							1	
- 1			gene	for thermostable			l		1			- 1
30	D288	879		polymerase I,			1		l A	F068748) cabi	1	- 1
_	~200			lete cds	0.	.56		3659692	liv I	F068743) sphingosine kinase lus musculus]		
31	Z150	27	ri.sap III DN	iens HLA class						43758) fibrinogen A-alpha-	0.	.77
			レ!	<u></u>	0.	56		1304141	ch.	ain	1 _	
- 1		- 1	Humai	n T-cell surface		ı			7		 0.	76
			intige	CD2 (TII)		i			- 1			- 1
32	<u>M</u> 143	62	nRNA	complete cds.	Δ.	., 1			(Y	11915) Tenascin-X [Bos		1
				Joinpiele eds.	0.	10	:	2462979	tau	rus)	0.5	.
-]		12	.mays	mRNA for		- 1			T		U.,	"
.		c	hlorop	hyll a/b-		- 1			1	ì		I
3	Z5080	<u>) </u>	inding	protein CP29	0.5	6		1004==	coll	agen alpha I(I) chain -		- 1
					<u> </u>	<u>~_</u>		109677	Imo	use >gi 50487	0.5	_ 1

_	<u> </u>	earest	Neighbor (BlastN	vs. Ger	nbank)	Nanear	NT :		
SE	Q]		į.			rearest	INCIE	hbor (BlastX vs. Non-Redunda	nt Proteins)
II	ACCE	SSION	DESCRIPTION	N	P VALU	1			
-				7		THE CLOS	1014	DESCRIPTION	P VAL
- [1							HYPOTHETICAL 86.0 KD	
j	1			- 1		ł		PROTEIN IN GLK 1-SRO9	
1	ł			- 1		1		INTERGENIC REGION	
1			S.cerevisiae	ĺ		1		>gi 83159 pir S19367	1
434	Z381		chromosome XIII			1		hypothetical protein YCL039)w -
	2.301	14	cosmid 9745	-	0.56	140372		yeast (Saccharomyces	1
1	1	Į,	G., b	Γ		1.03.72		cerevisiae)	0.35
1	1	- 1.	Escherichia coli D	NA		l		1	
435	AF0522	254	gyrase A (gyrA) go Partial cds	епе.				(AF038535)	
	- 552		egula pulligo 125		0.56	2724126		(AF038535) synaptotagmin V [Homo sapiens]	
1	1	S	mall subunit	'				(L-10100 Suprens)	0.12
	l		ibosomal RNA ge			j		1	1
1	j	m	itochondrial gene	iie,	٠.	!		1	1
	1	fc	r mitochondrial	- 1					
l	1	R	NA, partial	- 1	i		į	CYCLIN-DEPENDENT	1
436	AF0806	19 se	Quence	1	ا م		- 1	KINASE INHIBITOR I	1
			anio rerio mRNA	 -	0.56	3913223		p21/WAF1 [Felis catus]	
- 1		for	r protein tyrosine		- 1			į sas satusj	0.11
437	AJ00569	0 kir	nase	1,	0.56		- 1	(AF030962) unknown	1
- 1		Hu	man noggin	- 	7.30	2623830	!	Schistosoma mansoni]	7e-06
430		(N	OGGIN) gene,	- 1	- 1		1		70-00
438	U31202	Cor	nplete cds.	1 0	0.56	2075475		Z78411) F02D8.3	
					-	3875475		Caenorhabditis elegans]	le-06
439	Veren	Ovi	is sp. trichohyalin	1	1		L.		
~~	X51695	mR	NA, partial	0.	.56	3386622		AC004665) unknown protein	1
- 1		P			-	3300022	-11	Arabidopsis thaliana]	1e-10
- 1		Katt	us norvegicus	1	- 1		- 1		
_ [prot	ein tyrosine	1	- 1		CA	F072709) putative] i
40	U28938		sphatase D30	I	- 1		OX	idoreductase [Streptomyces] 1
		Timer	VA. complete cds	0.5	56	3293547	liv	idans]	
ı		Born	elia burgdorferi				7		1e-14
- 1		(secr	ion 57 of 70) of		- 1			·	
41 /	AE001171	the co	omplete genome	2 -	.		(A)	F016452) similar to the beta	
		1	ompicie genome	0.5	6	2315521	tra	asducin family	4
- 1		Caen	orhabditis		- 1		I(U	7951) contains similarity to	4e-16
			ns cosmid		- 1		AT	P/GTP-binding site motif	
2 A	F036685	C05B	10	0.56		(Class	(PS	:PS00017) [Caenorhabditis	
				0.50	'	1519671	eleg	ans]	6e-20
		Хепор	ous laevis		J				00-20
3 ,		vitello	genin gene Al		- 1		[(U4	1558) K02B2.3 gene	1
'	(01173	5' flan	king region	0.56	- 1	1118100	prod	luct [Caenorhabditis	
				3.50		1118102	eleg	ans]	2e-31
1			usculus DNA		1		l		
			2 protein.		ı		1	1]
<u> </u>	10911	omple	te cds	0.55	- 1	<none></none>	1	·. <none></none>	

F		Neare	st Neighbor (BlastN vs. (Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
5	SEQ					agnoti (Blasta vs. Non-Redundant	Proteins)	
F	ID	ACCESSIC		P VALUE	ACCESSION	DESCRIPTION	P VALU	
-	445	D20010	Rice mRNA EN117,				IF VALU	
<u> </u>	+43	D30010	partial sequence	0.55	<none></none>	<none></none>	-	
l l	ı		Escherichia coli			CHOINES	<none></none>	
14	146	Ite ioo.	phosphoprotein	1	ļ		1	
- -	 1	U51991	phosphatase	0.55	<none></none>	<none></none>	2102	
- 1	ı		Mouse T cell receptor			CHOILE	<none></none>	
1 4	47	Minnen	C-gamma-7.1 mRNA,		l			
 `	~/	M18858	3' end.	0.55	<none></none>	<none></none>	<none></none>	
i	ł		Homo sapiens T cell-			410112	KNUMES	
- 1	- 1		death associated		ĺ	i	ľ	
1 4	48	Mosala	protein gene,		•		1	
 -	"	U95218	complete cds	0.55	<none></none>	<none></none>	-NONTE:	
44	10	M14040	Human R-ras gene,			3.0.1.2	<none></none>	
 	" 	M14948	exon 1.	0.55	<none></none>	-\ <none></none>	ANONT	
			Human mRNA for			3.01.62	<none></none>	
45	. I	AB002353	KIAA0355 gene,	- 1				
177	"	AB002353	complete cds	0.55	<none></none>	<none></none>	-NONT	
-)	ſ		Homo sapiens				<none></none>	
1			(subclone 1_d6 from	j			1	
45	. 1	Y 0 1 600	PI H54) DNA	- 1		1	1 1	
1 43	┷╂╾	L81689	sequence	0.55	<none></none>	<none></none>	NOVE	
1	- 1		1			TATOME	<none></none>	
-1				ł			1 1	
1	1		Human myristoylated	1			1 1	
1			alanine-rich C-kinase	I		(AE001220) V-type ATPase,		
452	,	M68955	substrate (MACS)	ſ		subunit B (atpB-1) [Treponema]]	
- 132	+	14108933	gene, 5' end.	0.55	3322710	pallidum]	1 50 1	
ļ	1		D				5.0	
1	1		R.norvegicus mRNA	1		1	1	
453	1		(pJG116) with	1		extensin-like protein - maize		
1 733	╁		repetitive elements Synechocystis sp.	0.55	1076802	>gi 600118 mays]	5.0	
1	1		mntABC transporter]	3.0	
1	1		system: periplasmic-	i i				
1	1		oinding protein	1	•	1	.]	
ı				1]		
1	1		mntC), complete cds;	i		j l	1	
	1		mntA) gene,	- 1		hydrogen dehydrogenase (EC	ſ	
	1		omplete cds;	1		1.12.1.2) - Clostridium	1	
			nembrane protein	1		acetobutylicum >gi 557064	ı	
454	,		mntB) gene,			(U15277) hydrogenase I	1	
-,,,,	┝─┴		omplete cds.	0.55	2117632	[Clostridium acetobutylicum]	5.0	
	Ī		lasmodium berghei			g server accredity fleating	3.0	
			nerozoite surface	1	!		ł	
			rotein-1 gene. Omplete cds	1	j		1	
455	, ,			0.55				

V

SEO ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION P VALUE		Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
H. H. H. H. H. H. H. H.	SEO					Total Vs. Non-Redundant P	roteins)	
H.sapiens CpG DNA.	· ·		DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	PVALUE	
A56 Z64937 Clone 17g7, reverse read epg17g7.rtla D.55 A17298 MFS18 PROTEIN PRECURSOR 3.8							1	
A56 Z64937 read cpg17g.Tal.a. Macaca mulatta clone iris8.3 T-cell receptor alpha chain mRNA, partial cds. 0.55 310406 (L09212) tat protein [Simian immunodeficiency virus] virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] virus virus virus virus] 3.8 (L09212) tat protein [Simian immunodeficiency virus] virus viru							 	
Macaca mulatta clone inh83 T-cell receptor alpha chain mRNA, partial cds. Homo sapiens multidrug resistance protein 0.55 1585251 traB gene (Amycolatopsis methanolica) 2.8	1				•	MFS18 PROTEIN		
Macaca mulatta clone inh8a T -cell receptor alpha chain mRNA, partial cds. 0.55 310406 immunodeficiency virus] virus 3.8	456	Z64937		0.55	417298	PRECURSOR	3.8	
AFO U10914 U10914 AFO U10914 AFO U10914 AFO U10914 AFO U10914 AFO U10914 AFO U10914 AFO U10914 AFO U109	ŀ	Ī					1 3.3	
457 U10914 partial cds. D.55 310406 immunodeficiency virus virus 3.8 Homo sapiens multidrug resistance protein D.55 1585251 traB gene [Amycolatopsis methanolica] 2.8 Mouse Hox-3.1 gene and Hox-3.2-Hox-3.1 intergenic region. D.55 818849 [U25430] nucleotide pyrophosphatase precursor [Oryza sativa] 2.0 Plasmodium falciparum chromosome 2. section 32 of 73 of the complete sequence D.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete Sequence D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32 of 73 of the complete D.55 Section 32		l	•				1	
Homo sapiens multidrug resistance protein 0.55 1585251 traB gene [Amycolatopsis methanolica] 2.8	157	7710014				(L09212) tat protein [Simian	i	
Mouse Hox-3.1 gene and Hox-3.2-Hox-3.1 intergenic region. 0.55 1585251	437	010914		0.55	310406	immunodeficiency virus] virus]	3.8	
Mouse Hox-3.1 gene and Hox-3.2-Hox-3.1 intergenic region. 0.55 818849 (U25430) nucleotide pyrophosphatase precursor [Oryza sativa] 2.0								
Mouse Hox-3.1 gene and Hox-3.2-Hox-3.1 intergenic region. 0.55 818849 (U25430) nucleotide pyrophosphatase precursor [Oryza sativa] 2.0	150	A F022020						
Mouse Hox-3.1 gene and Hox-3.2 Hox-3.1 intergenic region. 0.55 818849 (U25430) nucleotide pyrophosphatase precursor [Oryza sativa] 2.0	438	AF022838	protein	0.55		methanolica)	2.8	
A59 M35603 Intergenic region. 0.55 818849			Mouse Hoy 2 Lane					
M35603 intergenic region. 0.55 818849 [Oryza sativa] 2.0	1 .		1	٠.		1	<u>[</u>	
Plasmodium falciparum chromosome 2 section 32 of 73 of the complete sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete Sequence 0.55 PROTEIN C2 Section 32 of 73 of the complete Sequence O.55 PROTEIN C2 Sequence 0.55 PROTEIN C2 Sequence 0.55 PROTEIN C2 Sequence 0.55 PROTEIN C2 Sequence 0.55 PROTEIN C2 Sequence 0.55 PROTEIN C2 PROTEIN C2 Sequence 0.55 PROTEIN C2 Sequence 0.55 PROTEIN C2	459	M35603		0.55	010040			
falciparum chromosome 2, section 32 of 73 of the complete sequence 0.55 137532 PROTEIN C2 >gi 74386 pir WZVZB6 59K HindIII-C protein - vaccinia virus (strain WR) 1.7 1.7	122	11133003		0.33	818849	[Oryza sativa]	2.0	
Chromosome 2, section 32 of 73 of the complete sequence 0.55 137532 137532 1.7								
Section 32 of 73 of the complete sequence 0.55 137532 Sgi 74386 pir WZVZB6 59K HindIII-C protein - vaccinia virus (strain WR) 1.7			· · ·			PROTEIN C2		
the complete sequence 0.55 137532 HindIII-C protein - vaccinia virus (strain WR) 1.7 Plasmodium falciparum chromosome 2, section 32 of 73 of the complete sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete sequence 0.55 137532 Virus (strain WR) 1.7 Human transcription factor (NFATc.b) mRNA. complete cds 0.55 3327144 [Homo sapiens] (AB014565) KIAA0665 protein (Homo sapiens) PROTEIN IN GLK1-SRO9 INTERGENIC REGION PROTEIN IN GLK1-SRO9 INTERGENIC REGION Sequence origin recognition complex large subunit (ORC1) gene, complete cds 0.55 140372 cerevisiae) 0.017 Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons (GCD	1							
AE001395 Sequence 0.55 137532 Virus (strain WR) 1.7			the complete					
Plasmodium falciparum chromosome 2, section 32 of 73 of the complete sequence 461 AE001395 sequence 0.55 137532 virus (strain WR) Human transcription factor (NFATc.b) mRNA, complete cds Saccharomyces cerevisiae origin recognition complex large subunit (ORC1) gene, complete cds 0.55 140372 (U46674) coded for by C. elegans cDNA yk27d9.5; coded for by C. elegans cDNA yk27d9.3; short region of weak homology to drosophilia	460	AE001395		0.55	137532	•	, ,	
chromosome 2, section 32 of 73 of the complete sequence 0.55 137532 PROTEIN C2 Section 32 of 73 of the complete sequence 0.55 137532 Section WR) 1.7			Plasmodium		201332	Viius (strain VVR)	1.7	
section 32 of 73 of the complete sequence 0.55 137532			falciparum					
the complete sequence 0.55 137532 HindIII-C protein - vaccinia virus (strain WR) 1.7 Human transcription factor (NFATc.b) (AB014565) KIAA0665 protein [Homo sapiens] (AB014565) KIAA0665 protein [Ho	ļ i		chromosome 2,			PROTEIN C2		
the complete sequence 0.55 137532 HindIII-C protein - vaccinia virus (strain WR) 1.7 Human transcription factor (NFATc.b) (AB014565) KIAA0665 protein [Homo sapiens] 0.096 Saccharomyces cerevisiae origin recognition complex large subunit (ORC1) gene, complete cds 0.55 140372 Cerevisiae) Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons (GCDH)	1 1		section 32 of 73 of	1		>gi 74386 pir WZVZB6 59K		
461 AE01395 sequence 0.55 137532 virus (strain WR) 1.7 Human transcription factor (NFATc.b) (AB014565) KIAA0665 protein [Homo sapiens] 0.096 Saccharomyces (Erevisiae origin recognition complex large subunit (ORC1) gene, complete cds 0.55 140372 cerevisiae) U34860 gene, complete cds 0.55 140372 cerevisiae) Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons (GCDH) gene, exons (GCDH) gene, exons (GCDH) gene, exons (GCDH) gene, exons (GCDH) gene, exons (AB014565) KIAA0665 protein [Homo sapiens] (Homo sapiens) (Homo sapiens) (Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons (GCD			•				*	
factor (NFATc.b) mRNA, complete cds U59736 factor (NFATc.b) mRNA, complete cds O.55 3327144 [Homo sapiens] HYPOTHETICAL 86.0 KD PROTEIN IN GLK1-SRO9 INTERGENIC REGION >gi 83159 pir S19367 hypothetical protein YCL039w - yeast (Saccharomyces cerevisiae) U34860 U34	461	AE001395	sequence	0.55	137532		1.7	
factor (NFATc.b) mRNA, complete cds U59736 factor (NFATc.b) mRNA, complete cds O.55 3327144 [Homo sapiens] HYPOTHETICAL 86.0 KD PROTEIN IN GLK1-SRO9 INTERGENIC REGION >gi 83159 pir S19367 hypothetical protein YCL039w - yeast (Saccharomyces cerevisiae) U34860 U34		·		f				
462 U59736 mRNA, complete cds 0.55 3327144 [Homo sapiens] 0.096 Saccharomyces PROTEIN IN GLK1-SR09 INTERGENIC REGION	1	ľ	•	- 1			Î	
Saccharomyces cerevisiae origin recognition complex large subunit (ORC1) gene, complete cds U34860 U	462		1				ſ	
Saccharomyces cerevisiae origin recognition complex large subunit (ORC1) gene, complete cds U34860 Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons 461 AFO12341 6 7 8 9 and 10 Saccharomyces INTERGENIC REGION >gi 83159 pir S19367 hypothetical protein YCL039w - yeast (Saccharomyces cerevisiae) (U46674) coded for by C. elegans cDNA yk27d9.5; coded for by C. elegans cDNA yk27d9.3; short region of weak homology to drosophilia	402	039736	mRNA, complete cds	0.55	3327144		0.096	
Saccharomyces cerevisiae origin recognition complex large subunit (ORC1) gene, complete cds Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons Saccharomyces INTERGENIC REGION >gil83159 pir S19367 hypothetical protein YCL039w yeast (Saccharomyces cerevisiae) (U46674) coded for by C. elegans cDNA yk27d9.5; coded for by C. elegans cDNA yk27d9.3; short region of weak homology to drosophilia	ŀ	4					1	
cerevisiae origin recognition complex large subunit (ORC1) gene, complete cds U34860		1	Saccharomyces					
recognition complex large subunit (ORC1) gene, complete cds U34860 Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons 464 AF012341 6 7 8 9 and 10			_		•			
large subunit (ORC1) gene, complete cds U34860 U3486			- 1	J		1	į	
463 U34860 gene, complete cds 0.55 140372 cerevisiae) 0.017 Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons (GCDH)	Į	ſ		ł		, ,		
Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons 464 AF012341 6.7.8.9 and 10	463		, ,,	0.55	140373	, ,		
Homo sapiens glutaryl-CoA dehydrogenase (GCDH) gene, exons 464 AF012341 6.7 8.9 and 10			gener, complete cus	0.55			0.017	
glutaryl-CoA dehydrogenase (GCDH) gene, exons 464 AF012341 6 7 8 9 and 10		[1	Homo sapiens				[
dehydrogenase (GCDH) gene, exons homology to drosophilia	1	ls	glutaryl-CoA				[
(GCDH) gene, exons homology to drosophilia	- 1			į			1	
461 AF012341 6 7 8 0 and 10 0 55 115551	- 1			[
	464	AF012341 6	5. 7. 8, 9, and 10	0.55			0.008	

SE	1		1		ghbor (BlastX vs. Non-Redundant	
-) IACCESSIC	1				1
1	TACCESSIC		P VALUE	ACCESSION	DESCRIPTION	P VALUE
-		HIV-1 isolate Q98-				I VALUE
1	l	CxA from Kenya,				
	<u> </u>	envelope		1	l l	1
1	İ	glycoprotein C2V3	1	1		1
465	4500400.	region (env) gene,	1	[1
40.	5 AF004891		0.54	<none></none>	<none></none>	<none></none>
466	Y 10159	D.discoideum				
+00	110139	racGAP gene Homo sapiens mRNA	0.54	<none></none>	<none></none>	<none></none>
		for B120, complete		Ì		
467	AB001895		0.54	4NONT-		
		Bovine gene for	0.54	<none></none>	<none></none>	<none></none>
		aspartyl protease		' .		
	1	NM1 exons 3 and 4 >			İ	
	I	:: lcl X12357 Bovine			1]
		aspartyl protease				1
		NM1 gene, exons 3				1 1
468	X12357	and 4.	0.54	<none></none>	<none></none>	<none></none>
		D- 1: 1				
	1	Borrelia burgdorferi	· .		ļ	1 1
469	AE001151	(section 37 of 70) of the complete genome	25.			1
,	742007131	H.sapiens mRNA for	0.54	<none></none>	<none></none>	<none></none>
	i	T cell receptor alpha		•	·	
470	X92052	chain	0.54	<none></none>	NONE	I I
		Mus musculus ileal	9.54	CHOILES	<none></none>	<none></none>
		lipid-binding protein	I		(U27698) calreticulin	1 1
471	U00938	gene, complete cds	0.54	1009712	[Arabidopsis thaliana]	4.9
	l				CASEIN KINASE II, ALPHA	7.2
	İ	,	j		CHAIN (CK II)	l f
	f	1	j		>gi 419938 pir A43297 casein	1 1
	1		1		kinase II (EC 2.7.1) alpha	1 1
		M thermafarminian			chain - Theileria parva	1 1
	4	M.thermoformicicum complete plasmid	ŀ		>gi 161871 (M92084) casein	·
472	X68367	pFZ1 DNA	0.54	125222	kinase II alpha subunit	į 1
		P. E.I DIAN	0.34	125272	[Theileria parva]	4.7
		H.sapiens CpG DNA.	ŀ		1	ļ i
ı		clone 44c4, reverse	1		(AJ131094) Xvent-1B protein	ļ <i>1</i>
173	Z61098	read cpg44c4.rt1a	0.54	4191274	[Xenopus laevis]	2.7
		Human gastric H,K-			[[.tonopus tucvis]	3.7
		ATPase catalytic	ļ		(Z70757) similar to serine	ŀ
., 1		subunit gene,	ļ		protease inhibitor	
74		complete cds.	0.54	3881648	[Caenorhabditis elegans]	3.7
.75		H. sapiens mRNA for			(D87963) ETF-related factor-1	
	730019	PRPL-2 protein	0.54	1648828	(ETFR-1)	2.1

Nearest Neighbor (BlastN vs. Genbank) SEQ ID ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION S.glaucescens genes strU, strX, strV and strW for 5'-hydroxystreptomycin pruduction and transport polypeptides 476 X89010 Homo sapiens mRNA for Hic-5, partial cds Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-monooxygenase	
ID ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION S.glaucescens genes strU, strX, strV and strW for 5'- hydroxystreptomycin pruduction and transport polypeptides 476 X89010 Polypeptides Homo sapiens mRNA for Hic-5, partial cds Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	D
S. glaucescens genes str U, str X, str V and str W for 5'- hydroxystreptomycin pruduction and transport polypeptides 476 X89010 Homo sapiens mRNA for Hic-5, partial cds Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	12111
str U, str X, str V and str W for 5'- hydroxystreptomycin pruduction and transport polypeptides 476 X89010 Homo sapiens mRNA for Hic-5, partial cds Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl- Str U, str X, str V and	P VALUE
strW for 5'- hydroxystreptomycin pruduction and transport polypeptides 476 X89010 Homo sapiens mRNA for Hic-5, partial cds Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl- (AF084524) cellular repress of E1A-stimulated genes CF [Mus musculus] ORF 1 [Streptomyces lavendulae]	
hydroxystreptomycin pruduction and transport polypeptides 0.54 3550345 (AF084524) cellular repress of E1A-stimulated genes CF [Mus musculus] Homo sapiens mRNA for Hic-5, partial cds 0.54 1097213 ORF 1 [Streptomyces lavendulae] Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	
pruduction and transport of E1A-stimulated genes CF [Mus musculus] Homo sapiens mRNA for Hic-5, partial cds Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	
transport of E1A-stimulated genes CF [Mus musculus] Homo sapiens mRNA for Hic-5, partial cds 0.54 1097213 ORF 1 [Streptomyces lavendulae] Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	ľ
476 X89010 polypeptides 0.54 3550345 [Mus musculus] Homo sapiens mRNA for Hic-5, partial cds 0.54 1097213 ORF 1 [Streptomyces lavendulae] Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	or
Homo sapiens mRNA for Hic-5, partial cds Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	ŒG
AB007836 for Hic-5, partial cds 0.54 1097213 lavendulae] Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	0.25
AB007836 for Hic-5, partial cds 0.54 1097213 lavendulae] Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	
Comamonas testosteroni TsaR (tsaR), toluenesulfonate methyl-	
testosteroni TsaR (tsaR), toluenesulfonate methyl-	0.15
testosteroni TsaR (tsaR), toluenesulfonate methyl-	!
(tsaR), toluenesulfonate methyl-]
toluenesulfonate methyl-	
methyl-	1 1
monooxygenase	1
oxygenase component	j
component (tsaB),	
toluenesulfonate zinc-	1
indepedent alcohol (Z96047) DY3.6	1 1
478 U32622 dehydrogenase 0.54 3875351 [Caenorhabditis elegans]	0.006
Arabidopsis thaliana	1 1
gene for beta-VPE, D61394 complete cds 0.53 cNONEs	
479 D61394 complete cds 0.53 <none> <none></none></none>	<none></none>
Arabidopsis thaliana	
gene for beta-VPE,	
480 D61394 complete eds	
M.capricolum DNA COMPLET COMPLETE COMPLICATION COMPLETE COMPLETE COMPLETE COMPLETE COMPLETE COMPLICATION COMPLETE COMPLETE COMPLETE COMPLETE COMPLETE COMPLICATION COMPLETE COMPLETE COMPLETE COMPLETE COMPLICATION COMPLETE COMPLETE COMPLETE COMPLETE COMPLETE COMPLETE COMPLETE COMPLET	<none></none>
481 733072 for CONTTIC MC007 0.52	
233072	<none></none>
Human	
phosphatidy linositol [17]	
(4,5)bisphosphate 5-	
phosphatase homolog	
482 U45975 mRNA. partial cds. 0.53 <none> <none></none></none>	<none></none>
S.cerevisiae S.cerevisiae	THORES
chromosome XIV	
reading frame ORF M130 antigen (cytosolic varia	nt
483 Z71324 YNL048w 0.53 2135586 1) - human	2.1
	
Listeria	
monocytogenes sec A (AF016415) No definition line	
484 L32090 gene, complete cds. 0.53 2291129 found [Caenorhabditis elegans	

	Neare	st Neighbor (BlastN v	s. Genbank)	Negrost N.	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ			J. Gollounk)	14cmest 14	eignbor (BlastX vs. Non-Redundan	t Proteins)		
ID	ACCESSIO			E ACCESSIO	N DESCRIPTION	P VAL		
		Mus musculus mR	NA			IP VAL		
485	20000	for HGT keratin.			(X96713) collagen [Globoder			
463	D86423	partial cds	0.53	1235974	pallida!			
	ľ	Mus musculus V				0.41		
486	3515050	kappa 21-6 gene,	í	1		1		
400	Y15969	partial	0.52	<none></none>	<none></none>	3,00		
		Mus musculus (clor			CHOILES	<non!< td=""></non!<>		
1 1		3F9) transcribed		l l	ł	- }		
		germline T cell		1		J		
1 1		receptor gamma cha	:-	1				
i i		(Ter-g) mRNA, VJ4		1	(Z67990) Similarity to Rat	1		
487	M27480	C4 region.	4		amiloride-sensitive sodium	1 .		
		Human (lambda)	0.52	3875542	channel beta-subunit	4.6		
1 1		DNA for						
i I		immunogloblin light	-	1	1	ı		
488	D87004	chain			(U37272) winged helix protein	1		
		Human DNA	0.52	1766073	CWH-1 [Gailus gallus]	3.5		
		sequence from	1					
		cosmid E75B8 on	1	ł		1		
- 1		chromosome 22.	1	1		ł		
		complete sequence		•	.	1		
489	Z99704	[Homo sapiens]	0.51	<none></none>	1			
- 1			0.51	CHOILES	<none></none>	<none></none>		
		Sambucus nigra lectir	ı]	f		1		
405		precursor mRNA,			7			
490	U76523	complete cds	0.51	<none></none>	<none></none>			
- 1		••			CHOILES	<none></none>		
1		Haemophilus	1 1		1	j		
- 1		influenzac Rd section			1	l		
491		110 of 163 of the				1		
7/1	U32795	complete genome	0.50	<none></none>	<none></none>	<none></none>		
		,	. Т		helicase homolog g10L protein -	ZIAOIAE>		
	l,	U	· . [African swine fever virus			
192		Human myoglobin			>gi 414091 (X72951) G10L 125			
		gene, exon 2.	0.49	478384	KDa protein	7.0		
- 1	1		j		(U97002) similar to	,		
J	- 1				Schizosaccharomyces pombe 4-			
J	1	luman mRNA for	1		nitrophenylphosphatase .			
- 1		SIAA0238 gene,	1		(PNPPASE) (SP:Q00472,			
.93		artial eds	024		NID:g5004) [Caenorhabditis	·		
	P	artial Cub	0.24	1938429	elegans}	2.5		
- 1	l _x	enopus laevis	ŀ					
- 1		nitotic	1					
- [hosphoprotein 90]]		
94 t		RNA. complete cds	0.23	NONE	1	}		
		P.O.C CG3	V.2.J	<none></none>	<none></none>	<none></none>		

	Nines					
-		est Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant	Proteins)
	EQ			1		
<u> </u>	D ACCESSI		P VALUE	ACCESSION	DESCRIPTION	P VALU
-		N.crassa				IF VALU
	i	mitochondrial small				
		(19S) rRNA and Cys	s- ·		(L05670) clustrin [Mus	J
4	95 J05254		0.23	192150	musculus]	5.1
- 1		Gene for glutamate			,	
	İ	dehydrogenase (EC			(L07867) invariant surface	1
49	6 X16399	1.4.1.4), put. bacteria			glycoprotein [Trypanosoma	ł
 	V 10399	Origin	0.23	790933	brucei]	0.030
-		Treponema pallidum	İ	Ì		
	i	section 67 of 87 of	i			1
49	7 AE00125	the complete genome	0.00		1	1
	1230123	the complete genome	0.22	· · <none></none>	<none></none>	<none></none>
		Homo sapiens		1		
ļ	ł	amyloid lambda light	!		·	
-	· I	chain variable region	1			j
49	AF026919	mRNA, partial cds	0.21	<none></none>	37035	
		D.melanogaster	5.27	CHOILES	<none></none>	<none></none>
499	Z27247	mRNA for defensin	0.21	<none></none>	<none></none>	1
		Candida albicans			KNONES	<none></none>
500	Y15608	UBI3 gene	0.21	<none></none>	<none></none>	<none></none>
501	1/00500	Human beta-tubulin				CHOILES
501	V00598	pseudogene.	0.21	<none></none>	<none></none>	<none></none>
1		A.thaliana		•		
	İ	microsatellite				
502	X79426	[repeated motif			1	}
100	1179420	A.caerulescens	0.21	<none></none>	<none></none>	<none></none>
1	I	mitochondrial genes				
1	1	for cytochrome b and			1	1 1
	i /	NADH	i		1	1
503	X75772	dehydrogenase 5	0.21	139626	PD 000000000000000000000000000000000000	i i
			<u> </u>	139020	PROTEIN TI PRECURSOR	7.8
1	ł	1	. 1		(AL031583) 1-	ļ į
	ł	1.	1		evidence=predicted by content;	
	i	1	ı		1-method=genefinder;084; 1-	
	1		i i		method_score=47.46; I-	ŀ
	I		1		evidence_end; 2-	
ì			J		evidence=predicted by match; 2-	· }
			- 1		match_accession=SWISS-	i
	4.	Serratia marcescens			PROT:P23792; 2-	1
504	A F020=2 (site specific	·		match_description=DISCONNE	
504	AF028736	recombinase	0.21	3645960	CTED PROTEIN.; 2-matc	4.6
		S	1			
505	X97545	S.cerevisiae OST5			(AF014940) No definition line	
	1577343	gene	0.21	2275631	found [Caenorhabditis elegans]	2.7

	Nearest	Neighbor (BlastN vs. C	ienbank)	Negrect Main	hhos/DimyV	
SEQ		Treignoor (Diagary vs. e	T T	realest Neig	hbor (BlastX vs. Non-Redundant P	roteins)
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DECORPTION	
			T VALUE	ACCESSION	DESCRIPTION	P VALUE
		Human prostate-			(1107012) (2015)	
1		specific antigen (PA)			(U97012) C04E6.2 gene	
506	M24543	gene, complete cds.	0.21	1938527	product [Caenorhabditis	1
		gone, complete cas.	0.21	1730327	elegans] RNA REPLICASE	2.7
		ĺ			POLYPROTEIN 2.7.7.48) -	1
1	ł	Mouse	Į.		Erysimum latent virus	ĺ
		thrombospondin	ŀ		>gi 3892232 (AF098523)	
	ľ	(THBS1) gene,	!		replicase protein [Erysimum	
507	M62470	complete cds.	0.21	548563	latent virusl	1
				340303	(L40584) polyprotein	2.1
		Homo sapiens cosmid			[Infectious pancreatic necrosis	
508	Y13544	Cı	0.21	1235710	virus]	2.0
		Chicken MHC B				2.0
		complex protein (C12				
		3) mRNA, complete			(AF090441) extracellular reelin	i .
509	M24193	cds.	0.21	3600102	[Gallus gallus]	0.52
		H.sapiens TFE3 gene,			(X83413) U88 [Human	0.52
510	X97161	exon 4,5 & 6	0.21	854065	herpesvirus 6]	0.30
					TRANSCRIPTION FACTOR	0.50
		R.norvegicus DNA			COUP 2 COUP-TFII - chicken	i
		sequence for	I		>gi 392817 (U00697) orphan	1 1
		LFB I/HNFI	*		receptor COUP-TFII [Gallus	
511	X67649	promoter	0.21	3913114	gallus]	0.004
ı			ł			
ı		Fugu rubripes growth				
512		hormone (GH) gene,			(AF030881) pol polyprotein	
312	U63807	complete cds	0.21	3510505	[Fugu rubripes]	3e-04
ľ		ĺ	1		NAM7 PROTEIN (NONSENSE	
			ļ		MEDIATED MRNA DECAY	
	l		į		PROTEIN 1) (UP-	
	j		ŀ		FRAMESHIFT SUPPRESSOR	ļ
ı	. [,	H.sapiens mRNA for	Į		1) factor NAM7 - yeast	
513		laminin alpha 5 chain	0.21	400250	(Saccharomyces cerevisiae)	· •
+		Mirounga leonina	0.21	400350	>gi 4023	le-07
		najor	1			
j		istocompatibility				ŀ
		complex class II	j			1
		DQA) gene, partial				
514		eds	0.20	∠NONE-	1027	
-		Fransmissible	0.20	<none></none>	<none></none>	<none></none>
		astroenteritis virus	ſ			
1		FI virion protein	ŀ			
515	~~~~	enes	0.20	<none></none>	NONE	NONE
		Prosophila virilis		3.10.112	<none></none>	<none></none>
1		imple DNA	ĺ			
16		equence (pDv-19)	0.20	<none></none>	<none></none>	NONE
		<u> </u>		11101112	<none></none>	<none></none>

	Neare	st Neighbor (BlastN vs. (Carbants	Normal			
SI	Q	SE THEIGHDON (DIASHN VS. (Jenbank)	Nearest Ne	ighbor (BlastX vs. Non-Redundan	t Proteins)	
	ACCESSIC	DESCRIPTION	P VALUE	i			
-					250ckii Holl	P VALUE	
İ		Homo sapiens biliary					
١,,	7	glycoprotein (BGP)	1	I	ľ		
51	7 M76741	gene, partial cds.	0.20	<none></none>	<none></none>	<none></none>	
		14.	1			KNONES	
1.		Mus musculus rae28			Į.	l	
51	8 D78515	gene, exon 1 and			Ţ		
1	D/8515	5'flanking region	0.20	<none></none>	<none></none>	<none></none>	
	I	Drosophila	[
	f	melanogaster RNA	1	•			
-	1	polymerase II second				1	
		largest subunit			J		
	1	upstream (DmRP		٠.	ì		
519	M62975	140) gene. exons 1-4.	0.20	<none></none>	1	1	
			0.20	CNONES	<none></none>	<none></none>	
	1	Chicken 78-kD	1			1 1	
1	i	glucose-regulated	į		j	ŀ	
520	M27260	protein, complete cds.	0.20	<none></none>	<none></none>	1,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
ı		Rice tungro			GVONES	<none></none>	
		bacilliform virus	Ĭ			1 1	
621	1 F00 4 1	Serdang strain,				1 1	
521	AF076470	complete genome	0.20	<none></none>	<none></none>	<none></none>	
1		Rice tungro				1410112	
1		bacilliform virus	j	•	1	1	
522	AF076470	Serdang strain,				1 8	
	111 070470	complete genome Human	0.20	<none></none>	<none></none>	<none></none>	
ı		cyclooxygenase-2	f				
		(hCox-2) gene,	1			1 .	
523	U04636	complete cds.	0.20	NONE		1	
		Plasmodium	0.20	<none></none>	<none></none>	<none></none>	
	}	falciparum	1		1	1	
]	chromosome 2,	1		j ,) !	
		section 67 of 73 of	1		·	1 1	
	1	the complete	1			1	
524	AE001430	sequence	0.20	<none></none>	<none></none>	1,10,7	
J					HYPOTHETICAL 15.5 KD	<none></none>	
	1		ſ		PROTEIN IN MOAE-RHLE	1 1	
Į		Mus musculus	1		INTERGENIC REGION	1 1	
Ī		phosphomannomutase	I		>gi 1787009 (AE000181) orf,] · [
525		Pmm2) mRNA,	J		hypothetical protein	ļ 1	
323	14 043314 0	complete cds	0.20	3025006	[Escherichia coli]	9.8	

	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Na	all was a second	
SEC	2		1	. rearest Ive	ghbor (BlastX vs. Non-Redundant F	roteins)
ID	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
\vdash	+					
526	U23144	Xenopus laevis F1Z F1-related nuclear orphan receptor variant (xFF1rAshor mRNA, complete cd	1)	3184402	(AB014477) period protein	
			0.20	3184402	[Chymomyza costata] PROBABLE MICROSOMAL	9.6
527	U14621	Paracentrotus lividus Pax-6 (suPax-6) mRNA, complete cds	1	465894	SIGNAL PEPTIDASE 23 KD SUBUNIT (SPC22/23) >gi 630688 pir S44854 K12H4.4 protein - Caenorhabditis elegans >gi 289708 (L14331) homology with signal peptidase; coded for by C. elegans cDNAs GenBank: M79661, M79662 and M79663;	
			0.20	403894	putative	7.7
528	AF030511	Actinobacillus pleuropneumoniae MRP ATPase homolog (mrp) gene, partial cds; ApxIVA var3 (apxIVA) gene, complete cds; and beta-galactosidase (lacZ) gene, partial cds	0.20	1175966	HYPOTHETICAL 45.3 KD PROTEIN IN THIS 5'REGION >gi 1084720 pir S56193 probable membrane protein YFL062w - yeast (Saccharomyces cerevisiae)	7.2
					glyoxal oxidase (EC 1.2.3)	7.2
529	AF070581	Homo sapiens clone 24540 mRNA sequence T.maritima pgK gene	0.20	542394	precursor - basidiomycete (Phanerochaete chrysosporium) >gi 1050302	5.8
530	X75437	for 3- phosphoglycerate kinase	0.20	825648	(Z34531) coproporphyrinogen oxidase [Homo sapiens]	5.8
531	•	Haemophilus influenzae Rd section l of 163 of the complete genome	0.20	3309593	(AF072878) ciliary outer arm	5.6
532	, ,	S.cerevisiae chromosome XI reading frame ORF YKL081 w	0.20		CARBON CATABOLITE DEREPRESSING PROTEIN KINASE >gi 1469803 (L78129) serine/threonine kinase [Candida albicans]	5.5

-	: Near	est Neighbor (BlastN vs	. Genbank)	Nearest Ne	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SE	Q				Silvor (BlastA vs. Non-Redundant F	roteins)		
ID.	ACCESSI		P VALUE	ACCESSION	DESCRIPTION	P VALU		
├─		Hordeum vulgare				IP VALU		
ł	1	limit dextrinase				+		
533		(HvLD99) gene,	1		(AF064077) adrenocorticotropic	1		
233	AF02272		0.20	3139154	hormone receptor [Sus scrofa]	1		
	i	Drosophila			instruction (Sus scrota)	4.3		
534	47.00.00	melanogaster cosmi	d	ł	(AC005623) putative argonaute			
234	AL02172		0.20	3885334	protein [Arabidopsis thaliana]			
	1	Brassica rapa mRNA			p [: Dubidopsis tilalialia]	2.6		
535	ABOLOTO	for SRK45, complete	e	1	(Z92824) B0413.4]		
333	AB012100	cds cds	0.20	4008334	[Caenorhabditis elevans]	1		
	i				60S RIBOSOMAL PROTEIN	1.5		
	1		Ì		L30B (RP29) cytosolic - yeast	· .		
	1				(Saccharomyces cerevisiae)			
	ł		1		>gi 171821 not determined)			
	j				[Saccharomyces cerevisiae]			
	l		1		>gi 1045254 cerevisiae]			
	Ī	H.sapiens HLTF gene	?		>gi 1323250 gnl PID e243708			
536	746606	for helicase-like	1		(Z72933) ORF YGR148c			
230	Z46606	transcription factor	0.20	132946	[Saccharomyces cerevisiae]	1.6		
537	V07102	H.sapiens mRNA for		-	DNA-REPAIR PROTEIN	1.5		
	X87193	2.19 gene	0.20	139820	XRCC1	1.5		
			1		HYPOTHETICAL 33.5 KD	1.5		
- 1			1 1		PROTEIN IN SEC53-ACT1			
- 1		1		•	INTERGENIC REGION			
- 1		Í			>gi 1084703 pir S56211			
		Clostridium			probable membrane protein			
- 1					YFL044c - yeast			
- 1		perfringens C beta 2	1		(Saccharomyces cerevisiae)			
38	L77965	toxin gene, complete			>gi 836711 gn PID d1009835	i		
	277903	cds	0.20	1175950	(D50617) YFL044C	1.4		
- 1		Chicken neural cell-	į					
- 1		adhesion molecule (N			1			
39	M15938			•	1			
+		CAM) gene, exon 18.	0.20	2133082	regulatory protein MSR1 - yeast HYPOTHETICAL 55.9 KD	1.1		
		1	ł		HYPOTHETICAL 55.9 KD			
ŀ			j		PROTEIN C56G2.1 IN	j		
		Solanum tuberosum	1		CHROMOSOME III	ŀ		
ı		mRNA for extensin-	1	•	>gi 726413 (U23177) C56G2.1	J		
0		like protein, partial	0.20		gene product [Caenorhabditis	ı		
_		nto protein, partial	0.20		elegans]	1.1		
1		ĺ	1		EPIDERMAL GROWTH			
ĺ	j		ſ		FACTOR PRECURSOR	ļ		
	}		I		precursor - mouse >gi 309210]		
1	X98108	A.thaliana psbP gene	222		(J00380) prepro-egf [Mus	1		
	17	anana psor gene	0.20	119227	musculus)	0.49		

AB01117	Homo sapiens mRN for KIAA0607 protein, partial cds H.sapiens ITIH1 ger (exon 22) and ITIH3	P VALUE	1	ghbor (BlastX vs. Non-Redundant P DESCRIPTION gene VGF protein - rat >gi 205690 (M60525) nerve growth factor inducible protein [Rattus norvegicus] >gi 205701 (M60522) nerve growth factor- inducible protein [Rattus	Proteins) P VALI
AB01117	Homo sapiens mRN for KIAA0607 protein, partial cds H.sapiens ITIH1 ger (exon 22) and ITIH3	JA 0.20	ACCESSION	gene VGF protein - rat >gi 205690 (M60525) nerve growth factor inducible protein [Rattus norvegicus] >gi 205701 (M60522) nerve growth factor-	
	for KIAA0607 protein, partial cds H.sapiens ITIH1 ger (exon 22) and ITIH2	JA 0.20		gene VGF protein - rat >gi 205690 (M60525) nerve growth factor inducible protein [Rattus norvegicus] >gi 205701 (M60522) nerve growth factor-	PVALI
	for KIAA0607 protein, partial cds H.sapiens ITIH1 ger (exon 22) and ITIH2	0.20	2143753	>gi 205690 (M60525) nerve growth factor inducible protein [Rattus norvegicus] >gi 205701 (M60522) nerve growth factor-	
	for KIAA0607 protein, partial cds H.sapiens ITIH1 ger (exon 22) and ITIH2	0.20	2143753	>gi 205690 (M60525) nerve growth factor inducible protein [Rattus norvegicus] >gi 205701 (M60522) nerve growth factor-	
	for KIAA0607 protein, partial cds H.sapiens ITIH1 ger (exon 22) and ITIH2	0.20	2143753	growth factor inducible protein [Rattus norvegicus] >gi 205701 (M60522) nerve growth factor-	
	for KIAA0607 protein, partial cds H.sapiens ITIH1 ger (exon 22) and ITIH2	0.20	2143753	[Rattus norvegicus] >gi 205701 (M60522) nerve growth factor-	
	for KIAA0607 protein, partial cds H.sapiens ITIH1 ger (exon 22) and ITIH2	0.20	2143753	(M60522) nerve growth factor-	
	for KIAA0607 protein, partial cds H.sapiens ITIH1 ger (exon 22) and ITIH2	0.20	2143753	inducible protein (Range	1
	H.sapiens ITIHI ger (exon 22) and ITIH		2143753	Inducible protein (Ramus	I
X75318	H.sapiens ITIHI ger (exon 22) and ITIH		2173733	Protein (Manus]
X75318	(exon 22) and ITIH			norvegicus] >gi 207651	0.39
X75318	(exon 22) and ITIH		1	RNA-binding protein rnpD -	
X75318		ne		Arabidopsis thaliana (fragment) >gi 510240 (X61108) RNA	
X/5318		3	٠.	binding protein (A01108) RNA	
		0.20	629557	binding protein [Arabidopsis thaliana]	
	Oncorhynchus mykis	is		thanala)	0.38
	mRNA for alpha 3	1 1		muf1 protein - human	
A B000274		1 1		>gil762953 (X86018) must	
AB008374	partial cds	0.20	1082610	[Homo saniens]	
		1 7		[same supremy	0.37
	I imulus malaaka	1 1		1	
		1 1	•	1	
U09809				(AJ012650) CP [Papaya	
	muctyA, complete cas	0.20	3882016	ringspot virus]	0.37
	Homo saniens mPNA	1 1	-		0.37
		1		(U93121) M-phase	
AB020671	1	0.00		phosphoprotein-1 [Homo	
	Phytophthora	0.20	2674350	sapiens]	0.18
	megasperma				
	mitochondrial				
	ORF152, complete	j			
	cds, cytochrome c	1			!
	oxidase subunit I	ď]	ł
	(cox1) gene,	1		i	
	complete cds,	1.		(133513) D	j
	cytochrome c oxidase	- 1		(U2351/) D1022.7]
	subunit II	0.20	746516	Caenornabditis elegans]	ł
			,40010	gijo208001 elegans]	0.043
		į.	j		
	mitochondrial	ł		1	ĺ
	ORF152, complete	- 1		l.	i
ľ	eds, cytochrome c	1		ľ	1
		1			- 1
]	1		1
C	complete cds.	1	la	123517) D1022 7	j
	ytochrome c oxidase	1			i
_U4457 s	ubunit II	0.20	746516	Actioniabultis elegans	ſ
	U09809 AB020671 L04457	Limulus polyphemus arginine kinase mRNA, complete cds Homo sapiens mRNA for KIAA0864 protein, partial cds Phytophthora megasperma mitochondrial ORF152, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit II Phytophthora megasperma mitochondrial ORF152, complete cds, cytochrome c oxidase subunit II (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase cytochrome c oxidase complete cds, cytochrome c oxidase complete cds, cytochrome c oxidase cytochrome c oxidase	Limulus polyphemus arginine kinase mRNA. complete cds. 0.20 Homo sapiens mRNA for KIAA0864 protein, partial cds 0.20 Phytophthora megasperma mitochondrial ORF152, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit II 0RF152, complete cds, cytochrome c oxidase subunit II (cox1) gene, complete cds, cytochrome c oxidase subunit II (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds. cytochrome c oxidase subunit I (cox1) gene, complete cds. cytochrome c oxidase	AB008374 type I collagen, partial cds 0.20 1082610 Limulus polyphemus arginine kinase mRNA, complete cds. 0.20 3882016 Homo sapiens mRNA for KIAA0864 protein, partial cds 0.20 2674350 Phytophthora megasperma mitochondrial ORF152, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome coxidase subunit II 0.20 746516 Phytophthora megasperma mitochondrial ORF152, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome cox	type I collagen, partial cds 0.20 1082610 [Homo sapiens] Limulus polyphemus arginine kinase mRNA, complete cds. 0.20 3882016 [Homo sapiens] Homo sapiens mRNA for KIAA0864 protein, partial cds 0.20 2674350 [U93121) M-phase phosphoprotein-1 [Homo sapiens] Phytophthora megasperma mitochondrial ORF152, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase subunit I (cox1) gene, complete cds, cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase cytochrome c oxidase

	-	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Ne	ghhor (Blood V V	
s	EQ				Theatest Ive	ghbor (BlastX vs. Non-Redundant F	Proteins)
	ID	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DEGGE TO THE STATE OF THE STATE	
			Cdk3r=cyclin-	T	T ACCESSION	DESCRIPTION	P VALI
- 1			dependent kinase 5	 	 		
	- 1		regulatory subunit	· I	1		
	- 1		p35 [mice, brain,	1		ł	1
			129/SvJ, C57BL/6,	1			
- 1	- 1		Genomic/mRNA,			(ł
54	49	S82819	5528 nt]	0.20	3413870	(AB007923) KIAA0454 protein	·Į
1			Streptomyces griseus	1	3413670	[Homo sapiens]	0.020
- 1	- 1	,	DNA for	1	ł	1	
	- [serine/threonine	l .		(1/2015 t) TOTAL	1
	- 1	•	protein kinases.	ĺ	<u> </u>	(U29154) T07F12.2 gene	i
55	0	D31792	complete cds	0.20	861405	product [Caenorhabditis	
	ı		Homo sapiens	0.20	801403	elegans]	0.019
	1		butyrophilin (BT3.2)			(AE04005 ()	
1	- 1		gene, exons 5-10, and	i		(AF040954) putative protein	
55	1	<u>U</u> 97499	complete cds	0.20	2773341	phosphatase I nuclear targeting	
1	1			5.50	2773341	subunit [Rattus norvegicus]	0.008
1	- [Rattus norvegicus			1	
1	1		nonmuscle myosin				
1	1		heavy chain-A			(781130) 1	
552	<u>'</u>	U31463	mRNA, complete cds.	0.20	3880111	(Z81130) predicted using Genefinder	
1	1				5000111	Genetinder	0.002
1			Bacteriophage P22				
1	1		right operon, orf 48,	i		1	
1	1		replication genes 18	- 1			
1			and 12, nin region	1		1	
ļ			genes, ninG	1		1	
l	1		phosphatase, late	1		1	
1	1		control gene 23, orf	ſ		1	
i	1		60, complete cds, late	i i		(U42436) C49H3.3 gene	
553	Ι,		control region, start	j		product [Caenorhabditis	
333	 -	X78401 c	of lysis gene 13	0.20	1123087	elegans]	4e-0-1
	i	,	Nocardia				46-04
				ŧ	•	·	
			actamdurans pcbAB nd pcbC genes for	1	1	1	
		la a	ing hence Reues tot	1			
			lpha-aminoadipyl-L-	1		PUTATIVE ENDONUCLEASE	
			ysteinyl-D-valine	1		C1F12.06C yeast	l
			one initial	1		(Schizosaccharomyces pombe)	1
554	Y		openicillin N Inthase	l ·		>gi 1217980 (Z69944) unknown	í
	^			0.20	1723511	[Schizosaccharomyces pombe]	4e-09
ł			epidermidis genes	1		3, 50001	37
Į			piY', epiY, epiA,	1	1		1
555	x		oiB. epiC, epiD, oiQ. epiP	1	le le	(Z73424) C44B9.1	i
		-2300 [e]	iQ. Epir	0.20	3874927	Caenorhabditis elegans]	3e-10

-	Ne.	arest	Neighbor (BlastN	vs. Ger	bank)	Nearest Nai	ghhor (Planty	
SEC	2		ľ			- Addition (VC)	ghbor (BlastX vs. Non-Redundant I	Proteins)
ID	ACCES:	SION	DESCRIPTIO	N	VALUE	ACCESSION		
				-		ACCESSION		P VAL
1 .				_ -			(Z80220) similar to nucleotide	7
1				- 1		1	binding protein; cDNA EST	1
1	I			- 1			EMBL:M75897 comes from thi	s
1	i			- 1		1	gene; cDNA EST	1
l	1	- 1		1		İ	EMBL:M89054 comes from this	s
1	l		Epizootic	- 1		[gene; cDNA EST	ĺ
		ŀ	haemorrhagic disea	ase		i	EMBL:D26713 comes from this	İ
ارمما		ľ	virus gene segment	6		1	gene; cDNA EST	1
556	X5900	0 1	for NSI		0.20	38797 <i>55</i>	EMBL:D26718 comes from this	1
557	3 40 0		Human keratin 1			3017133	gene; cDNA	8e-16
557	M9877		gene, complete cds	- 1	0.20	1086900	(U41278) contains similarity to	
			Mus musculus			1000300	G beta repeats	2e-30
558	A F0.44		granzyme K gene,					-
220	AF01144	6 c	omplete cds		0.19	<none></none>	310375	
		١,				2.0.1.22	<none></none>	<none< td=""></none<>
ı		I,N	lacaca mulatta clos	ne	ı		1	
- 1		I ^M	1MUI.5 FRG1-like	:	l			
- 1		ĮP:	seudogene, exons 7	7	I		1	
559	AF074708	- 1	nd 8. partial	- 1	ı		1	
	7.1-074708		quence		0.19	<none></none>	<none></none>	
560	X13287		edicago sativa				GAOIVES	<none></none>
-	7113207	100	dulin-25 gene cerevisiae	4).19	<none></none>	<none></none>	
- 1			romosome X	4			- CATORIES	<none></none>
				1				
561	Z49509		nding frame ORF R009c	1 .	1			
	4.7507		vine DNA for	1 0	.19	<none></none>	<none></none>	NONT
.			staglandin	1	- 1			NONE>
- 1		F2:	alpha receptor.	1	1	Ì		
62	D89041	par	tial cds	1 .			1	
		Stre	ptococcus	<u> </u>	19	<none></none>	<none></none>	NONE>
ı	•		varius DNA for	1	- 1			.10112
63	D29644	dex	tranase	1				
T		Hei	icobacter pylori,	0.	19	<none></none>	<none></none>	NONE>
- 1		strai	in J99 section 22		- 1			
		of 1	32 of the		i	1		ł
4 A	E001461	com	plete genome	0.1	.	310315		•
		Hon	no sapiens	0.1	"	<none></none>	<none></none>	IONE>
-		galac	tocerebrosidase		- 1	1		
		(GA	LC) gene, exon		1	į.	. 1	
5 I	_38559	17.		0.19	, l	NONE	·	ł
1		R.pro	owazekii	0.1		<none></none>	<none> <n< td=""><td>ONE></td></n<></none>	ONE>
1		geno	mic DNA		J	1		
. [1	fragn	nent (clone		- 1		1	- 1
5 2		A405		0.19	. 1	NONE	l	1
	L		- /	0.19		<none></none>	<none> <no< td=""><td>ONE</td></no<></none>	ONE

	Neare	est Neighbor (BlastN vs. (Genhank)	Name		-
9	EQ		Jenoank)	Nearest Neig	ghbor (BlastX vs. Non-Redundan	t Proteins)
- 1	D ACCESSIO	ON DESCRIPTION	P VALUE		DESCRIPTION	
_				1	DESCRIPTION -	P VALUE
- 1		Tetrahymena		 		
l	1	thermophila]		ı
		telomerase	ł		1	ł
- 1	1	component p80			1	
50	57 U25641	mRNA, complete cds	0.19	<none></none>		
ŀ		Human mRNA for	1	CHOILES	<none></none>	<none></none>
- 1	j	KIAA0345 gene,	i		1	
56	8 AB002343	complete cds	0.19	<none></none>		
ł			- 5:17	CHOINES	<none></none>	<none></none>
	1	Erwinia carotovora				
1	_ i	gene for pectate lyase		٠.		
56	9 D10064	III. complete cds	0.19	<none></none>	1	1
-		Homo sapiens clone	0.17	GIONES	<none></none>	<none></none>
1	ł	MIF118 A4A10			i	1 1
		hypoxanthine			ł	}
	i i	phosphoribosyltransfe	j			
	ł	rase (hprt) 130 kb	İ			1
1	1 .	deletion mutant				1 1
1	1	mRNA, partial cds,	1		1	1 1
		contains human Alu	I			1 1
570	U31734	element	0.19	<none></none>	2703-	
		Plasmodium	<u> </u>	CHOINES	<none></none>	<none></none>
1		falciparum	1		1	1 1
i	1	chromosome 2,	1			1
1	1	section 23 of 73 of	1			1 1
1		the complete	1		1	1 1
571	AE001386	sequence	0.19	<none></none>	NONE	1 1
i	1	Homo sapiens		41OHD2	<none></none>	<none></none>
ĺ	ì	hydroxymethylbilane	ı		ĺ	1
1	1	synthase gene,	Į.			1
572	M95623	complete cds.	0.19	<none></none>	<none></none>	NONE
		(CC+15)			THORES	<none></none>
	[(GC*IS)=vitamin D-	ł		·] . [
		binding protein/group	į.			i i
		specific component	į.			i i
		[human, peripheral	İ]]
		blood leukocytes,	i]]
573		Genomic, 794 nt,	ŀ	. 1		1 1
2,2		segment 4 of 9]	0.19	<none></none>	<none></none>	<none></none>
574	.	H. sapiens NRGN				
		gene, exon 1	0.19	<none></none>	<none></none>	<none></none>
ľ		Homo sapiens	1			
ı		preakpoint cluster	i			i
575		egion BCRder14	. 1			
<u> </u>	712-044//3 S	equence	0.19	<none></none>	<none></none>	<none></none>

	Neares	Neighbor (BlastN vs. C	SE1-S	T 30		
SEC		Treignooi (Blastia VS. C	Jenbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant P	roteins)
ID	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
L		Human mRNA for	- 1	T		I VALUE
		KIAA0335 gene,				
576	AB002333	complete cds	0.19	<none></none>	<none></none>	<none></none>
		Macaca mulatta pit- 1/GHF-1 transcription factor			probable membrane protein	
577	U535 6 6	mRNA. complete cds	0.19	1078068	YLR311c - yeast	0.2
578	1172664	Human t(11;14)(q13;q32) breakpoint junction		. ,	COAT PROTEIN (CAPSID PROTEIN) virus >gi 58901 (X62133) CyMV coat protein	9.2
1-3/°	U73664	sequence Heterophyilaea	0.19	116734	gene product	8.8
579	AF004054	pustulata rps 16 gene, chloroplast gene, partial intron sequence	0.19	1928991	(U92815) heat shock protein 70	
			0.15	1928991	precursor [Citrullus lanatus] HYPOTHETICAL ATP-	8.7
580	Z 27081	Caenorhabditis elegans cosmid M01A8, complete sequence [Caenorhabditis elegans]	0.19	2496247	BINDING PROTEIN MJ0625 >gi 2128413 pir A64378 hypothetical protein MJ0625 - Methanococcus jannaschii >gi 1591336 (U67510) M. jannaschii predicted coding	
		S.cerevisiae	0.19	2496247	region MJ0625	8.6
581		chromosome IV reading frame ORF YDL097c	0.19	1174405	TYROSINE-PROTEIN	
		Small round structured virus genomic RNA, 3 terminal sequence containing ORF2 and	0.19	1174425	KINASE SPK-1	6.7
582		ORF3	0.19	971318	(Z48053) putative protein [Bovine herpesvirus 1]	5.1

	Near	est Neighbor (BlastN vs.	Genbank)	Nearest Neis	hbor (BlastX vs. Non-Redundant l	Oma a di N
s	EQ				Jase Va. Non-Redundant	roteins)
	D ACCESSI	ON DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	Byaren
<u> </u>		каняюща епиорпа			Deckii non	P VALUE
		DNA 16S ribosoma		1		
1	1	RNA > ::	1		1	1
	i	dbj D88002 D88002			İ	1
- 1	1 .	Ralstonia eutropha	1		·	ļ.
1.		DNA for 16S	1	l	.}	ļ
	- 1	ribosomal RNA > ::		1	ļ	1
	}	dbj D88003 D88003	1	ł		
-	1	Ralstonia eutropha		1		1
		DNA for 16S	ł	1		1
-	ļ	ribosomal RNA > ::		ļ.		1 1
		dbj D88004 D88004	ļ	i '·		1 1
1		Ralstonia eutropha		1]	İ
		DNA for 16S	1		(AE100657) N. 1.5	J i
58	3 D88000	ribosomal RNA	0.19	3800952	(AF100657) No definition line	
1		Methanococcus		2000932	found [Caenorhabditis elegans]	5.1
ł	ł	jannaschii section 4	1.		(AJ005586) MYB-related	1
1	ľ	of 150 of the	1 1		transcription factor	ł I
58	4 U67462	complete genome	0.19	3183617	[Antirrhinum majus]	
		Gallus domesticus		3103017	[[Antiffinum majus]	4.0
		microsatellite DNA	1 1		(U93074) voltage-gated sodium	1. 1
58	L23906	marker.	0.19	1947094	channel homolog BdNa1	1 20 1
1					Thanker Holliolog Bulva i	3.9
1		1			GLUCOSE-6-PHOSPHATE]
1	ı	_	1 1		ISOMERASE (GPI)	
1	i	Helicobacter pylori,	1 1		ISOMERASE) (PHI)	i
1	1	strain J99 section 23	i i		>gi 2118333 pir [148073 glucose	
		of 132 of the	l . f		phosphate isomerase - Chinese	1
586	AE001462	complete genome	0.19	1730177	hamster >oil987046 griseus1	3.9
ł	1				(282256) cDNA EST	- 3.7
ĺ		Ì			yk251g7.3 comes from this	i
ĺ	1	n			gene; cDNA EST yk251g7.5	I
l	1	P.putida catBC			comes from this gene; cDNA	1
		operon encoding			EST EMBL:D68223 comes	- 1
1	I	cis,cis-muconate	•		from this gene; cDNA EST	ļ
	I	lactonizing enzyme 1	ł		EMBL:C12737 comes from this.	ı
	I	and muconolactone	i		gene; cDNA EST yk389c8.5	- 1
587	MIGAGO	isomerase genes,			comes from this gene; cDNA	- 1
387	M19460	complete cds.	0.19	3873843	E	3.9
		Tetrahymena australis				
	Í	telomerase RNA	1		ļ	- 1
588	U22349	gene, complete			(AF049922) PGP169-12	1
200	022349	sequence	0.19	4105782	[Petunia x hybrida]	3.2

	Neare	st Neighbor (BlastN v	. Genbank)	Nearest Ma	ighhor (PleasY Y	
SEC	2			- Garest Ive	ighbor (BlastX vs. Non-Redundant l	Proteins)
ID	ACCESSIO	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VAL
	 	Homo sapiens volta				I VAL
1		operated calcium	ige]		+
	I	channel, alpha-1		i		
1	I	subunit mRNA,	1			
589	L27745	complete cds.	0.19	27/2024	(AC004450) unknown protein	1
1		Canis familiaris	0.19	3763926	[Arabidopsis thaliana]	3.0
1		synapsin I gene,	1	1	(45040405)	7
590	AF049588	partial cds	0.19	4104931	(AF042196) auxin response	I
1 1				4104931	factor 8 [Arabidopsis thaliana] PRE-NECK APPENDAGE	3.0
j		1 .	1	i	PROTEIN (LATE PROTEIN	
1 1				l · ·	GP12) >gi 75856 pir WMBP22	1
				1	gene 12 protein - phage phi-29	İ
					>gi 215330 (M14782) pre-neck	1
- 1		Staphylococcus	1 1		appendage protein	1
		aureus plasmid pS19	.1		[Bacteriophage phi-29]	
591	X06627	sequence	1		>gi 225367 prf 1301270G gene	
		M.musculus gene for	0.19	137927	12 [Bacteriophage phi-29]	2.3
1		kallikrein-binding	1 1	•		
592	X61597	protein	0.19	2002024	(AE000675) cobalamin	
			1 3.12	2982874	synthesis related protein CobW	1.7
1		Dictyostelium	l 1			
- 1		discoideum protein	1 1		1	
1		synthesis elongation	1		1	
593	AF016242	factor 1-alpha (tef2)	1		PUTATIVE RNA-DIRECTED	
323	AF016242	gene, partial cds Venezuelan equine	0.19	133659	RNA POLYMERASE	1.4
		encephalitis virus	1			1.4
- 1		strain 1327			1	
		polyprotein gene,	- 1		1	
		partial cds > ::	į.		1	
	į.	gb AF004460 AF004	ſ	:	1	
- 1		460 Venezuelan	- 1		· · · · · ·	
- 1		equine encephalitis	· 1		1	
- 1		virus strain 1385	- 1	ı		
94 4		olyprotein gene,	1	ŀ	(U25968) early embryogenesis	
"	AF004447	artial cds	0.19	4096173	protein [Oryza sativa]	,,
	ı	Juman alassia (DI an	1		INHIBIN BETA B CHAIN	1.3
		Iuman elastin (ELN) ene, exon 1, clones	J	1	PRECURSOR inhibin precursor	
- 1	la Li	ELC-5 and HELC-	İ	[.	- bovine >gi 563753 (U16241)	ı
95	J04821 6		0.10	į t	betaB inhibin/activin precursor	i
		omo sapiens histone	0.19	11/0523	Bos taurus]	1.3
1		eacetylase 3	}	. F	PROBABLE TRANSPORT	
		IDAC3) gene.	1	F	PROTEIN CY21C12.11	- 1
6 A		omplete cds	0.19	3024881	gi 2078066 gnl P1D e315171	j
				3024001	Z95210) betP	0.83

	Nea	rest Neighbor (E	BlastN vs. C	Genbank)	Nearest Na	sighbor (DLV	
S	EQ				. readest 146	eighbor (BlastX vs. Non-Redundant	Proteins)
	D ACCESS	ION DESCR	IPTION	P VALU	ACCESSION		P VALUE
\vdash						FERREDOXIN-DEPENDENT	
	i i			1		GLUTAMATE SYNTHASE I	
ł	ı	ł			I	(FD-GOGAT)	
1	1	- 1		ĺ	1	>gi 2126524 pir S60228]
- 1		- [1	glutamate synthase (ferredoxin)	,
- 1	1	D.melanoga	ostar			(EC 1.4.7.1) gltB -	1
		calcium-act				Synechocystis sp. (PCC 6803)	
_ 59	7 M6905			0.10		>gi 515938 (X80485) glutamate	.
		Dictyosteliu		0.19	1707984	synthase	0.80
J	1	firmibasis p			I		
- 1	- I	Dfp1, comp			1.		1
59	8 AF07627	9 plasmid sequ		0.19	453986	(U00008) yejA [Escherichia	1 1
- 1				0.15.	433980	coli]	0.79
- 1	1	Mouse MCN	IP gene		ł		
	1	for C-type na	atriuretic		1 .	(AE001002)	ľ
		peptide, com	plete cds		ļ	(AE001092) acetyl-CoA synthetase (acs-1)	1 1
599	D28873	(exoni, exon	2)	0.19	2650444	[Archaeoglobus fulgidus]	1. 1
	1					[Achieoglobus fulgidus]	0.63
	ı	Oxytricha no]
600	U06071	macronuclear				complement control protein	1
1000	000071	gene, comple	te cds.	0.19	1584024	[Botryllus schlosseri]	0.48
601	L54057	Homo sapien				(AL022374) putative ABC	0.46
	234037	mRNA, partic	al cds.	0.19	3036883	transporter	0.46
1	1	P.lividius cDi	VA So-	Ï		(AC004877) sco-spondin-mucin-	0.40
602	X89806	COLL2alpha		[like; similar to P98167 uncertain	1
		Archaeoglobu	gene	0.19	3638957	[Homo sapiens]	0.41
1	1	fulgidus section	on 3 of				
1		172 of the con	npiete	ł	•	1	ſ
603	AE001104	genome		0.19	2215102	(Y11739) transcription factor	1
1		Rattus norvegi	CUS	- U.13	2315192	[Homo sapiens]	0.35
1		microsatellite		1		D-MeAsp	
604	U54501	sequence DOM	lco22	0.19	228951	receptor:ISOTYPE=epsilon3	
]]					220731	[Mus musculus]	0.32
			1			(AF096371) contains similarity	ł
		Human		1		to Rattus norvegicus cyclin G-	j
605	V74466	papillomavirus	type	1		associated kinase (SW:P97874)	ſ
505	X74468	15 genomic DN	IA	0.19	3695390	[Arabidopsis thaliana]	0.28
		U	2001			(AJ002527) glucitol-6-	V.20
606	U20285	Human Gps1 (C				phosphate dehydrogenase	ſ
	020203	mRNA. comple Human gene for	te cds	0.19	2582659	[Clostridium beijerinckii]	0.27
j		interleukin 3 rec		[
- 1		alpha subunit, e	YOU	- 1		(AF008596) alpha1,3-	ĺ
607	D49408	10		0.19		fucosyltransferase [Helicobacter	I
				0.17	2522368	pylori].	0.16

 		st Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant P	modeline ()
SE	Q				- Compact vs. Holl-Redundant P	roteins)
П	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU
-		Homo sapiens	-			
	1	pituitary specific	1			— —
	1	homeodomain protei		i	1	
1	ł	(PROPI) gene, exon				
608	8 AF041141	3 and complete cds	0.19	37403	(X03541) trk gene product (aa 1	1
1		Discopyge ommata		37403	641) [Homo sapiens]	0.091
1		Ca2+ channel alpha	ı 🛭	1		ĺ
		subunit gene	1	ľ		
609	L12531	sequence.	0.19	3618274	(AJ223219) hypothetical protein	0.069
ł		Yellow fever virus			- September 2015	0.069
1	1	clone HONG9	1	, ,		
610	AF052445	polyprotein gene.			(U15928) KH-domain putative	
<u> </u>	71.032443	complete cds B.anthracis sap gene	0.19	1932822	RNA binding protein	0.001
]	1	encoding S-layer]]			
611	Z36946	protein	0.19	172044	(L06487) ZIPI protein	
		Homo sapiens full	0.19	173241	[Saccharomyces cerevisiae]	2e-04
		length insert cDNA			(AC005400) b	
612	AF087984	clone YW29A12	0.19	3786014	(AC005499) hypothetical	
		Archaeoglobus		3700014	protein [Arabidopsis thaliana]	1e-06
	1	fulgidus section 97 of			1	
£12		172 of the complete	1		(AF060248) unknown	į
613	AE001010	genome	0.19	3135493	[Arabidopsis thaliana]	7e-08
			1			70 00
	}	Trichosporon	Į.			
		cutaneum carbamoyl	i		1	1
		phosphate synthetase	I		(1741000	Ĩ
		large subunit (argA)	- 1		(U41278) F33G12.3 gene	
614	L08965	gene, partial cds.	0.19	1086901	product [Caenorhabditis	ĺ
		Rattus norvegicus		1000701	elegans]	2e-08
		A2b-adenosine	1			1
		receptor mRNA,	1	,	(AE000773) acetoin utilization	
615	M91466	complete cds.	0.19	2984320	protein [Aquifex aeolicus]	6e-09
ı					protein [Aquifex aeolicus] (AL032643) similar to	00-07
			i		Uncharacterized protein family	
ľ	- 1		ſ		UPF0034, Double-stranded	J
- 1	1	j	J		RNA binding motif; cDNA EST	1
	İ		1		yk489b3.5 comes from this	
- 1	ļs	S.lividans groEL2	1		gene; cDNA EST yk439g7.5	
16	320	ene	0.19		comes from this gene	
					[Caenorhabditis elegans] (U97016) similar to drosophila	7e-10
ı		Schizosaccharomyces	1		Rici gene product ribosomal	. 1
., [ombe sed2 (sed2)	1		protein L4 (YML4)	- 1
17	U12539 g	ene, complete cds.	0.19		(NID:g459259)	1

	: **	Near	rest Neighbor (BlastN vs.	Genbank)	Nearest No.	the (DL) V	
	SEC	2			iveriest IASI	ghbor (BlastX vs. Non-Redundant I	Proteins)
	ID		ION DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
	 					(U97016) similar to drosophila	T. VALUE
		ł	Schizosaccharomyce	ಜ		Ric1 gene product ribosomal	 -
	618	111252	pombe scd2 (scd2)			protein L4 (YML4)	
	1010	U1253	gene, complete cds.	0.19	1938549	. (NID:g459259)	9e-15
			Human DNA sequence from cosmid U25D11,			EMBL:D32434 comes from this gene; cDNA EST EMBL:D33710 comes from this gene; cDNA EST EMBL:D34467 comes from this gene; cDNA EST EMBL:D35005 comes from this gene; cDNA EST EMBL:D37535 comes from this gene; >gi 3878710 gnl PID e1348373 EST EMBL:D33710 comes from this gene; cDNA EST EMBL:D34467 comes from this gene; cDNA EST EMBL:D34467 comes from this gene; cDNA EST	S
			between markers	l [gene; cDNA EST	
l	619	Z68327	DXS366 and DXS87			EMBL:D37535 comes from this	
۲	<u> </u>	200327	on chromosome X. Dictyostelium	0.19	3875774	gene;	6e-15
	620	U66525	discoideum ORFveg114 mRNA, complete cds	0.19	3540281	(AF056116) All-1 related	
l					3340201	protein [Fugu rubripes]	2e-17
	521	U25830	Newcastle disease virus isolate Herts/33 matrix protein mRNA, complete cds Mus musculus strain	0.19	2228750	(U93868) RNA polymerase III subunit [Homo sapiens]	le-18
6	22	U89407	BALB/c delta- aminolevulinic acid dehydratase (Lv) mRNA, partial cds	0.19	1825764	(U88314) C46H11.11 gene product [Caenorhabditis elegans]	3e-25
6:	23	AF095598	Bison bison athabascae microsatellite BBJ 2	0.18	<none></none>		NONE>
62	24	AF064260	Strongylocentrotus purpuratus SRC8 mRNA, complete cds	0.18	<none></none>		NONE>

<u> </u>	Nea	rest Neighbor (I	BlastN vs. C	enbank)	Nearest N	leighbor (Blact V N.	
SEC	ર				1	leighbor (BlastX vs. Non-Redundar	t Proteins)
ID.	ACCESS	ION DESCR	IPTION	P VALUE	ACCESSIC	DESCRIPTION	P VAI
	1	Arabidops	ic thalians				
l	1		ha mRNA,				
625	U6953	complete c		0.10			
		Bovine DN	A for	0.18	<none></none>	<none></none>	NON
	1	prostagland	1		j		
	ł	F2alpha rec	eptor,		1	†	ĺ
626	D89041		·	0.18	<none></none>		
		Dictyostelii			CIVOIVES	<none></none>	<non< td=""></non<>
		discoideum			1		
627	1 ,6	Glu-GAA g	ene, clone		1		1
627	M24571			81.0	<none></none>	<none></none>	
	l	D.melanoga	ster ovo			CHOINES	<non< td=""></non<>
	ł	gene require	d for		1		
628	X59772	female germ			i	1	
	1537112	developmen Plasmodium		0.18	<none></none>	<none></none>	<non!< td=""></non!<>
		falciparum I					TON
		SEQUENCI	NGIN			1	1
1		PROGRESS	***				1
		from contig				1	
629	AL010209	complete seq		0.18	NONE		1
		Methanococo	us	0.18	<none></none>	<none></none>	<none< td=""></none<>
- 1		jannaschii se	ction 117	ł		1	
		of 150 of the		ı		inosital 1 4 5 -: 1	1
530	U67575	complete gen	ome	0.18	111839	inositol 1,4,5-triphosphate receptor 2 - rat	1
Í		Caenorhabdit				receptor 2 - rat	8.5
531	U28730	elegans cosmi	d	1		(AE000232) orf, hypothetical	
-	028730	K10B2		0.18	1787604	protein [Escherichia coli]	8.3
		L.lactis pepFI				(AF079110) glycosomal malate	0.3
32	X99798	pepF2 genes	~			dehydrogenase [Trypanosoma	1
				81.0	3406624	[brucei]	8.1
1				1	•	PROBABLE NUCLEAR	
		Danio rerio ba	nd 4.1-	1		ANTIGEN herpesvirus 1 (strain	1
		like protein 4 (nbl4)	1		Kaplan) >gi 334072 (M34651)	
33	AF025306	mRNA, compl	ete cds	0.18	465445	ORF-3 protein [Pseudorabies	
					.05 ++5	virus)	7.9
		Mus musculus		ł	•		
4 /	A EOSOSS I	lipoxygenase (a		1		(Z81368) hypothetical protein	
- - '	AF059251	mRNA, comple	te cds	0.18	1655667	Rv2393	6.6
5	7700	G.domesticus C	ľ			3-methyl-2-oxobutanoate	6.6
+	222003	protein mRNA.		0.18	481864	dehydrogenase	6.6
1		Homo sapiens n	DNA	l			0.0
1		for KIAA0514	2017	ı		1	
5 A		protein, complet	e cds	81.0	2024) 22	(Z81464) predicted using	
			- 203	,.10	3874158	Genefinder	6.4

35.13	1,40	st Neighbor (BlastN vs	. Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	ı			1.1000.031.146	gnbor (BlastX vs. Non-Redundant P	roteins)	
ΙD	ACCESSIO	DESCRIPTION	P VALUE				
		Caenorhabditis	PVALUE	ACCESSION	DESCRIPTION	P VAL	
		elegans cosmid					
	j	C07A4, complete	1	1			
	1	sequence	ł	1		l	
	İ	[Caenorhabditis		j	(AJ011681) retinoblastoma-	ļ	
637	Z 78536	elegans]	0.18	2700101	related protein [Chenopodium	Ì	
			0.18	3702121	[rubrum]	6.4	
		1.		1	(Z81094) Weak similarity to 65		
- 1		}	j	j	KDA heat shock protein		
		Methanococcus		l	(TR:G602231), cDNA EST	,	
- 1		jannaschii section 72	:1	i .	EMBL:D71705 comes from this gene; cDNA EST		
638	***	of 150 of the		٠.			
038	U67530	complete genome	0.18	3877946	EMBL:D74382 comes from this gene [Caenorhabditis elegans]		
1					(Z69634) cDNA EST	6.3	
		, ,	1 1		EMBL:D71510 comes from this		
			1 1		gene; cDNA EST		
- 1		1	1 1		EMBL:C08449 comes from this		
			1		gene; cDNA EST yk266b12.3		
.		Influenza	j i		comes from this gene; cDNA		
- 1		A/Duck/England/1/62	1		EST yk266b12.5 comes from		
		(H4N6) nucleoprotein			this gene; cDNA EST		
539	M63781	mRNA, complete cds.	0.18	2000 444	yk461h7.3 comes from this		
T			0.18	3873663	gene; cDNA	6.2	
		Oryctolagus					
•	•	cuniculus integrin			1		
- 1	į	beta-8 subunit	[1		
		mRNA, complete cds.	j		1		
- 1		> :: gb I44828 I44828	ł	•			
10		Sequence 3 from	1		major allergen OLE17 -		
~		patent US 5635601	0.18	1362129	common olive	e 0	
u l		D.melanogaster Rop	T		(AB011527) MEGFI [Rattus	5.8	
+	440/417	gene	O.18	3449286	norvegicus]	4.8	
	,	Homo sapiens beta-			[PROTEIN-PII]	7.0	
		rrestin 2 mRNA.	- 1		URIDYLYLTRANSFERASE		
2 A		complete cds		4	vinelandii >gi 39257 (X59610)	ĺ	
\top		ompiete cus	0.18	548353	uridylyl transferase	3.7	
1		Danio rerio	1				
	1.	untingtin (HD)	i	[1	potential IGF binding protein	ı	
3 A		RNA, complete cds	0.18	241050	chickens, Peptide Partial, 77 aa,	j	
				241058 s	segment 2 of 3]	3.6	

	Ne	arest Neighbor (Bl	astN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	EQ				carest 1	reign	oor (BlastX vs. Non-Redundant	Proteins)
-	D ACCES.	SION DESCRI	PTION	PVALUE	ACCESSIO	NC	DESCRIPTION	P VAL
				ļ			(Z68314) predicted using	1
1	1						Genefinder; cDNA EST	+
	1				1		EMBL:M75775 comes from the	nis
1	ľ				ł		gene; cDNA EST	-
İ	1	1	- 1		ĺ		EMBL:M89255 comes from th	is
ı	- 1		- 1		1	·	gene; cDNA EST	- f
1	į.	. Homo sapien	s mRNA		*		EMBL:M89127 comes from th	is
ı	1	for KIAA090	2				gene; cDNA EST	1
64	4 AB0207			81.0	3075550	ŀ	EMBL:T00141 comes from this	s
		HIV-1 isolate	patient	0.10	3875570		gene; cDNA EST EMBL:T	2.1
	1	3 country US	A pol		٠.	- 1		
	.1	polyprotein (p	ool)					1
645	AF0968	33 gene, partial o	ds	81.0	3250696	- 1.	(AT 00440.6)	1
	I	Pyrocoelia mi	yako		2520090		AL024486) putative protein	1.7
	. [(clone pB-Pm	L41)					
646	1 20000	luciferase mR	NA,			1	AC003974) umles-	1
U+0	L39928	complete cds		0.18	2914702	Jr.	AC003974) unknown protein Arabidopsis thaliana]	1
	ł	Human	- 1			- 1	. Habidopsis thananaj	0.73
	1	carcinoembryo	.	j				l
	1	nonspecific	піс	1		- 1		
	ł	crossreacting a		1		i		
	į .	(CEA; NCA) g	nugen	İ				
647	M17082	exons I and 2.	ene,	0.10		R	EGULATORY PROTEIN	
		2.		0.18	1351833	A	BAA	0.72
	i	1		1		RI	VA-binding protein rnpD -	
	i	H.sapiens ITIH	gene	- 1		Ar	abidopsis thaliana (fragment)	-
اا		(exon 22) and I	ГІНЗ	1		>g	i 510240 (X61108) RNA	
48	X75318	gene		0.18	629557	Dir	iding protein [Arabidopsis	
ı					029337	tha	liana]	0.41
J		Mus musculus	- 1					
- 1	•	apoptosis associa	ated			-		j
		tyrosine kinase	1	- 1		1:		
19	A FOLLOOS	(AATYK) mRN	A;	1		CKO	03332) nuclear antigen 2	Ï
	AF011908	complete cds		0.18	330442	(En	stein-Barr virus]	l
- [1	ł			ASI	ARTYL-IRNA	5e-04
- [1			SYN	NTHETASE aspartate	Į
-		Simian	- 1	1		tRN	A ligase (EC 6.1.1.12) -	į
		immunodeficienc	,	- 1		Esch	nerichia coli coli]	- 1
		virus SI VagmVEI		- 1		>gi	1736513 gnl PID d1016401	i
1		envelope protein	`-	ł		(D90	0829) AspartatetRNA	
2	U04004	gene, partial cds.		110		ligas	e (EC 6.1.1.12)	
J		Xenopus laevis	- - '	0.18	135102	[Esc	herichia coli]	6e-11
	- 1	RanGTPase	- 1	1				
上		activating protein	0	.18	005714	(X91	258) pid:e198503	j
				 -	995714	II Sacc	charomyces cerevisiae)	2e-13

1	Near	est Neighbor (BlastN vs.	Genbank)	Nearest Nai	ghbor (Blood V 27	
SEQ	2			Treatest Iver	ghbor (BlastX vs. Non-Redundant I	roteins)
ID	ACCESSI	ON DESCRIPTION	P VALUE	ACCESSION		PVALU
	 				(Z66511) similar to ribokinase;	1
1	ł		ł		cDNA EST EMBL:D69553	
ĺ	ı	1		ł	comes from this gene; cDNA	1
	l	1	1		EST EMBL:D65938 comes	1
	j	j			from this gene; cDNA EST	i
	1	B.oleracea gene for S	.	ı	yk280h9.3 comes from this	1
	ł	receptor kinase-like	"		gene; cDNA EST yk280h9.5	ł
652	Z18921	protein	0.0		comes from this gene; cDNA	
		S.cerevisiae STA2	0.18	3875535	EST yk223d11.3 come	le-19
653	M60650	gene, complete cds.	1 016			<u> </u>
		gone, complete cus.	0.16	<none></none>	<none></none>	<none< td=""></none<>
- 1		Eucalyptus globulus	J			
ı		NADP-isocitrate				
		dehydrogenase	1 1		1	
		(EgICDH) mRNA,	1 1		(AF057298) ornithine	
654	U80912	complete cds	0.16	2766170	decarboxylase antizyme 2 [Mus	
			0.10	3766172	musculus]	4.2
		Sambucus nigra	!!!			
		ribosome inactivating	1		1	
		protein precursor	1		<u> </u>	
655	AF012899	mRNA, complete cds	0.16	76749	hypothetical protein 4 - fowl	
				70749	adenovirus 1	4.0
1		Arabidopsis thaliana	. 1		1	
ı		cellulose synthase	ŀ		1	
		catalytic subunit (Ath-	ľ		1	
556	4 F00=+= :	B) mRNA, complete	j		(AF055904) unknown	
230	AF027174	cds	0.16	3044086	[Myxococcus xanthus]	0.00
- 1		Church			tycoods xandids]	0.60
- 1		Glycine max sucrose	j]	!
57	AF030231	synthase (SS) mRNA,	1		1	1
	ru 030231	complete cds	0.078	<none></none>	<none></none>	NONE>
- 1				•	cell wall glycoprotein, 75K.	
ı	1	Woodchuck hepatitis	1		precursor - diatom	
ı		virus (WHV),	1	İ	(Cylindrotheca fusiformis)	- 1
- 1		complete genome,	1		>gi 515363 (X80394) P75K	
8		clone WHV 59.	0.070		gene product [Cylindrotheca	i
\top		Ovine adenovirus	0.072	1076190	fusiformis]	6.3
1		Va2 protein gene,	1	ſ		
		ONA polymerase	1	1	. [j
-		gene, terminal protein	ł	İ	1	İ
-		ene and 52.55 kDa	1			ł
		protein gene, partial	1			- 1
9 1		ds gene: partial	0.072	(AF061244) unknown	į
			V.U/2	3511143	Agrocybe aegerita]	6.2

	Neares	t Neighbor (BlastN vs.	Genbank)	Nearest Neig	thbor (BlastX vs. Non-Redundant P	
SE			1	T.Carest 14c1g	moor (Blastix Vs. Non-Redundant P	roteins)
II	- 1		P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Caenorhabditis				II VALUE
	1	elegans cosmid				
		Y44A6B, complete		ł		
1	· I	sequence	1	Ī		
		[Caenorhabditis			·	1
660	AL021491	elegans]	0.070	<none></none>	<none></none>	<none></none>
		X.laevis Xotch				
66	M33874	protein mRNA,	0.070		(Y09076) RAD3	1
100	14153874	complete cds. Mus musculus	0.070	1654096	[Schizosaccharomyces pombe]	0.23
- 1		ZAN75 mRNA for		i		
		zinc finger protein,	1	,	MITOCHONDRIAL	
662	AB012725	complete cds	0.069	1350800	MITOCHONDRIAL	
		Caenorhabditis	0.000	1330800	RIBOSOMAL PROTEIN S5	2.0
ı		elegans cosmid	1			i I
- 1		Y44A6B, complete				i i
1		sequence				1
1		[Caenorhabditis				
663	AL021491	elegans]	0.068	<none></none>	<none></none>	<none></none>
1		H.sapiens CpG DNA,			(
		clone lel, reverse	1 1		(U55376) F16H11.2 gene	
664	Z60318	read cpglel.rla.	0.068	1280134	product [Caenorhabditis	
			0.000	1200134	elegans]	2.6
ĺ	ł				COA:3-KETOACID-	1
1			1		COENZYME A	
1	ļ		1		TRANSFERASE PRECURSOR	İ
1	}				EMBL:Z14816 comes from this	Ĭ
1]		l.		gene; cDNA EST	- 1
1	I .		1		EMBL:Z14946 comes from this	1
1	ĺ	S.cerevisiae	1		gene; cDNA EST	
1	1	chromosome II]		EMBL:D69746 comes from this	i i
1		reading frame ORF		. •	gene; cDNA EST yk219b6.3	ŧ
665		YBR104w	0.068	2402000	comes from this gene; cDNA	
		Streptomyces lividans	0.000	2493000	ES	0.68
		rpsP, trmD, rplS,	į.			- 1
	l li	sipW, sipX, sipY,	1.		1	1
]]:	sipZ, mutT genes and	j			1
i i		4 open reading			(X96713) collagen [Globodera	
666	Z86111	frames	0.068	1235974	pallida]	4e-04
	'					
			ŀ		(AC005164) match to ESTs	1
					AA731149 (NID:g2140138),	1
! I	را .	Anthonomus grandis			AA731908 (NID:g2752719),	ĺ
		itellogenin gene]		AA287837 (NID:g1933519),	1
667		VTG), complete cds.	0.068		AA262811 (NID:g1898382),	15.50
			3.330	3242730	and AA825820 (NID:g2899132)	le-59

	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Ne	ighbor (Blast Viva No. 1)	
SI	EQ.			.veurest 14e	ighbor (BlastX vs. Non-Redundant	Proteins)
Į.	D ACCESSIO	DN DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU
-						1 VALU
66	38 3424161	Rat tachykinin (PPT				
 "	8 M34161	gene, exons 5 and 6.	0.067	<none></none>	<none></none>	<none:< td=""></none:<>
	i	Aspersillus minum min				CINOINE
- 1		Aspergillus niger zin finger protein (creA)		i	1	1
66	9 L03811				.[ı
	203011	gene, complete cds. Human fibrinogen	0.067	<none></none>	<none></none>	<none></none>
Ţ	1	beta chain gene.	Í			13.10.112
1	I	1	İ	·	·	1
1	j	complete mRNA. >	1			1
	1	gb I47706 I47706	1			İ
670	M64002	Sequence 3 from		٠.		
10/	M64983	patent US 5639940	0.067	<none></none>	<none></none>	<none></none>
1		Nicotiana tabacum				CHOINES
1.	1	Mg chelatase subunit			}	1
671	AFOLAGE	(ChiH) mRNA,			1	1
10/	AF014051	partial eds	0.067	<none></none>	<none></none>	<none></none>
672	Y07540	ļ.,	1 1		glycoprotein GP330, renal - rat	CITOITES
0/2	10/340	H.sapiens sil gene	0.067	92331	(fragments)	7.5
1	ď				25 KD OOKINETE SURFACE	, · · · · · · · · · · · · · · · · · · ·
i	1		1		ANTIGEN PRECURSOR	
1	1	Rattus norvegicus			(PRS25) >gi 320962 pir A44966	
j	1	mRNA for 3'(2'),5'-			25k ookinete surface antigen	
673	A 10000 47	bisphosphate			precursor - Plasmodium	
0/3	AJ000347	nucleotidase	0.067	129238	reichenowi reichenowi]	7.4
i	1				hypothetical protein MJ0750 -	
ĺ	1				Methanococcus jannaschii	1
674	T 10070	Squid sodium channel			>gi 1592304 (U67521)	Į.
0/4	L19979	mRNA, complete cds.	0.067	2128473	ferredoxin-type protein	1.5
		Yeast tRNA-Glu(3)			ypo protein	
675	Vocaza	gene and flanking	- 1		(X15081) MURF2 protein (AA	
0/3	X08050	regions	0.067	1334398	1-348)	0.65
]				HYPOTHETICAL 51.6 KD	0.03
'	'	• .	ľ		PROTEIN CY49.14C	· [
İ	!		Į		>gi 1370241 gn1 PID e247089	:
•		Human mRNA for	j		(Z73966) hypothetical protein	-
676	V171.5	IgM heavy chain			Rv2075c [Mycobacterium	- 1
0/6	X17115	complete sequence	0.067	1731331	tuberculosis]	0.51
ľ	[ALPHA-2C-1 ADRENERGIC	0.31
ſ	1		. 1		RECEPTOR human >gi 178194	i
ſ	J.		1		(J03853) kidney alpha-2-	l
		Iomo sapiens	I		adrenergic receptor [Homo	ſ
ĺ		incoupling protein 3			sapiens] >gi 1628638 (U72648)	- 1
677		UCP3) gene, exon 1	<u> </u>		alpha2-C4-adrenergic receptor	
677	AF032871 a	nd partial exon 2	0.067	112900	[Homo sapiens]	0.50
					(supicing)	U.JU

	Nearest	Neighbor (BlastN vs. C	Genbank)	Nearest Neigh	ibor (BlastX vs. Non-Redundant Pr	roteins)
SEQ						
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		†	 		DYNAMIN 3 (DYNAMIN,	1 VALUE
	 		 		TESTICULAR) rat	
	1	Mouse class II MHC	I		>gi 391872 gnl PID d1003668	1
j		E-beta 2 (d) gene	1		(D14076) testicular dynamin	İ
678	X05319	exon 3	0.067	585074	[Rattus norvegicus]	3e-04
		Candida albicans			[[control cont	30 01
1		CaSLN1 gene,		,	(AC003007) Unknown gene	i
679	AB006362	complete cds	0.067	3417296	product (partial) [Homo sapiens]	9e-56
1						
1		African horse				1
1		sickness virus capsid		ĺ		
(00		VP3 (L3) mRNA,		! ' '		
680	AF021236	complete cds	0.066	<none></none>	<none></none>	<none></none>
1 1	1	Helicobacter pylori, strain J99 section 68		1	1	
		of 132 of the	1			
681	AE001507	complete genome	0.066	AIONE	NO.T.	
001	AL001307	Caenorhabditis	0.000	<none></none>	<none></none>	<none></none>
		elegans cosmid		İ		
682	AF039717	R13H8	0.066	<none></none>	<none></none>	<none></none>
· · · ·		Syncerus caffer		4.0.0.	NOTE:	CHOILS
		isolate Queen				
		Elizabeth Mweya 14				1
		mitochondrial DNA				
683	AF029027	control region	0.066	<none></none>	<none></none>	<none></none>
		Homo sapiens full		,		
604	. =======	length insert cDNA			(X97203) C1 protein [Beet curly	
684	AF087967	clone YU51G05	0.066	2982476	top virus]	9.5
!		ŀ			i .	
		Baboon endogenous			(I 27848) POIT tomain	
[[virus proviral long			(L37868) POU-domain	
685		terminal repeat DNA.	0.066	972767	transcription factor [Homo	7.3
-555	-,0207	terminar repeat DIVA.	0.000	7/2/0/	sapiens]	/.3
]	Lycopersicon	,		1	
		esculentum class I	ļ		1	
i	1:	knotted-like	į		(AC002131) Strong similarity to	
	l:	homeodomain protein	- 1		extensin-like protein gb Z34465	
1		(LeT6) mRNA,	j		from Zea mays. [Arabidopsis	
686	AF000141	complete cds	0.066	3157926	thaliana]	5.6
		Bensingtonia sp.			· ·	
ľ		OK255 gene for 18S	ĺ			- 1
- 1		rRNA > ::	ļ		[j
		dbj AB001747 AB00	ļ			J
J		1747 Bensingtonia	j `	•	(AF070064) cap 'n' collar	
407		sp. OK259 gene for			isoform C (Drosophila	
687	AB001746	18S rRNA	0.066	3859889	melanogaster]	0.38

7. 7.	Nearest	Neighbor (BlastN vs. (Genbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant P	roteins)
SEQ				5		Toterns,
l ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	D. V. A. I. W.
		Helicobacter pylori,	1	cerodion	DESCRIPTION	P VALUE
		strain J99 section 22	 			-
1		of 132 of the				
688	AE001461	complete genome	0.065	<none></none>	<none></none>	<none></none>
		Chicken erythroid				TATOM
		transport proteins c1	1		1.	
689	M30821	and c2	0.065	<none></none>	<none></none>	<none></none>
1		Homo sapiens gene				
		for osteonidogen,	Ì		1	
690	AB009802	intron 3	0.065	<none></none>	<none></none>	<none></none>
		Homo sapiens full		ĺ.	1 .	
691	AF086062	length insert cDNA clone YZ06B11	0.065	,		
1071	A1 080002	Human mRNA for	0.065	<none></none>	<none> SIGNAL SEQUENCE</none>	<none></none>
l ,		KIAA0371 gene,			BINDING PROTEIN binding	1
692	AB002369	complete cds	0.065	2500884	protein [Synechococcus sp.]	5.5
				2300004	protein (Syncendededs sp.)	3.5
		Cyclopodia sp. large				
1 1		subunit ribosomal		,		1
		RNA gene,				
		mitochondrial gene for mitochondrial				
		RNAs, partial				<u> </u>
		sequence > ::			1	
		gb AF086866 AF086				i I
i		866 Penicillidia sp.				! !
ŀ		large subunit				
i		ribosomal RNA gene,				
ı		mitochondrial gene	ł			1
		for mitochondrial	ı		ĺ	
		RNAs, partial			(AB012957) probable glycosyl	1
693	AF086864	sequence	0.065	3721684	transferase [Vibrio cholerae]	5.5
		Danis Die T				
		Bacteriophage BK5-T ORF410, 3' end pf	1			
		cds, 20 ORFs.				Ī
ļ		repressor protein, and				
		Cro repressor protein	1			
		genes, complete cds,			PEPTIDASE T	
		ORF70' gene, 5' end			(AMINOTRIPEPTIDASE)	
694		of cds.	0.065	1172067	intluenzae Rd]	3.2
		Ciona intestinalis	j	ł	'	
- 1		MyoD-family protein	İ			
ا برر		CiMDFa) mRNA,			(AL035353) contains EST	1
695	U80079	complete cds	0.065	4218110	gb:F15281	2.5

	. 1	Vearest	Neighbor (BlastN vs	Genbank)	Neprose N		
s	EQ				1 Actual Me	eighbor (BlastX vs. Non-Redundar	t Proteins)
	-	10122	DESCRIPTION	P VALU	E ACCESSIO	N DESCRIPTION	P VALUE
-			7.7				IF VALUE
ı	- 1		Homo sapiens mRN	A			
16	96 AR02		for KIAA0911	ŀ		MINOR CAPSID PROTEIN	7.3
<u> </u>	ABU2	20718	protein, complete co	is 0.065	1722734	>gi 1020192 type 23]	
	1		Zea mays copia-like			- B-1-020132 type 25]	1.9
i			retrotransposon Stl-			İ	l i
69	7 4 500	0	14 leader region,			(U89278) polyhomeotic 2	Į i
100	7 AF08	2137	partial sequence	0.065	1877501	homolog [Homo sapiens]	
			R.norvegicus ZnBP			g (=zomo sapiens)	1.1
69	8 X64	nen	gene for zinc binding	g			i
 	3 A041	033	protein	0.065	464963	TRYPSIN PRECURSOR	0.36
			M	1	•	- I LOOKOK	0.36
	i		Mus musculus	1		1	1 1
ŀ			butyrophilin (BTN)	1	j	1	1 1
699	9 U670		gene, promoter region		1	hypothetical protein YPL263c	. 1 1
 0	00/0	/03	and complete cds	0.065	2132252	yeast	
		[D	1	1	(AF041105) organic anion	3e-10
700	M648		Rat matrin F/G	1	i	transporter protein 3 [Rattus	
	171040	02	mRNA, complete cds Yeast (S.ccrevisiae)	0.065	3420183	norvegicus]	4e-19
1	1		ranscriptional	1			1 40-13
ł			ectivator of amino	ĺ	i	1] [
1	1		cid-biosynthetic		<u> </u>	1	1
1	1		genes (GCN4) gene,	1			1 1
701	K0220	25	omplete cds.	000	. 0	1	1 1
	1		Maize mRNA for a	0.064	<none></none>	<none></none>	<none></none>
	ł		igh mobility group				
702	X5828	2	rotein	0.044			
			lomo sapiens	0.064	<none></none>	<none></none>	<none></none>
1	ł		subclone 1_f3 from	1			
		P	I H69) DNA	f			i I
703	AC0015		quence	0.064	.	İ	1 1
			omo sapiens	0.004	<none></none>	<none></none>	<none></none>
1			RA3B region				
704	AF02346	- ·	quence	0.064	More	i	1
			enorhabditis	0.004	<none></none>	<none></none>	<none></none>
1 1		,	gans cosmid				
705	U50307		3Н9.	0.064	-NONE.	1	
		Sti	reptococcus crista	0.007	<none></none>	<none></none>	<none></none>
			npA gene, partial	j			
1 1			s, pulative	Į		(D83650) TDD	
i I			nesin/ABC	ł		(D83659) TPR protein pombe]	1
i I		tra	nsport system	i		>gi 2894282 gnl PID e1251103	ł
		pro	tein (scbA) gene,	[(AL021838) pre-mrna splicing	j
706	U46542		nplete cds	0.064	1209391	factor. [Schizosaccharomyces	1
					1207391	pombe]	9.2
70-			usticana mRNA	1		(H60315) MC004P (24.11	
707	X57564	lor	neurral peroxidase	0.064	1492037	(U60315) MC094R [Molluscum contagiosum virus subtype 1]	
			··		-1°C -0	comagiosum virus subtype []	6.9

	Near	est Neighbor (BlastN v	s. Genbank)	Negrec Mai	obb (D)	
SEC	2			Treatest Mei	ghbor (BlastX vs. Non-Redundant P	roteins)
D	7		P VALUE	ACCESSION	DESCRIPTION	Dyver
		Human alpha-2-				PVALL
1		macroglobulin				
í	1	receptor/lipoprotein	n			1
1	1	receptor protein		1	rab15B protein - wheat	ł
700		(A2MR/LRP) gene		İ	>gi 21853 (X62476) rab protein	1
708	U06986	exons 39-41.	0.064	100800	[Triticum aestivum]	
1	I	Human CpG island			(U88325) suppressor of	5.3
709	205	sequence, clone		i	cytokine signalling-1 (Mus	l
709	D85773	Q28B8	0.064	2245382	musculus]	1
	l				(AF096370) contains similarity	5.3
1	i .		1	}	to a C. elegans hypothetical	
ł	ĺ	1 • •	1		protein F44G4.1 (GB:Z49910)	
i	i		1		and several yeast hypothetical	
l	i	Apis mellifera			proteins such as 35.1 KD	
l	ľ	ligustica complete	1		protein in NAM8-GAR1	
710		mitochondrial	1		intergenic region (SP:P38805)	
710	L06178	genome	0.064	3695379	[Arabidopsis thaliana]	
					HYPOTHETICAL 70.5 KD	3.2
		j	1		PROTEIN IN AGP3-DAK3	
			1 1		INTERGENIC REGION	•
	-	1	1 1		>gi 1084712 pir S56201	
ľ		ĺ	1 1		probable membrane protein	
- 1		L	1		YFL054c - yeast	
		Triticum aestivum	1 1		(Saccharomyces cerevisiae)	
711	3/1/01/0	mRNA for beta-	1 1		>gi 836701 gn1 PID d1009825	
	Y16242	amylase	0.064	1175958	(D50617) YFL054C	٠.
		Homo sapiens			(C50017) 11 E054C	3.1
		(subclone 2_a2 from	1 1		1	
712	I 6 . 770	P1 H25) DNA	1 1		(AE001391) phosphatase (acid	
712	L81779	sequence	0.064	3845169	phosphatase family)	0.01
- 1		C.reinhardtii psb1			, , , , , , , , , , , , , , , , , , , ,	0.81
		mRNA for OEE1	i			
.		protein of].		(M94535) ATPase	
j	İ	photosystem [I	j		[Saccharomyces cerevisiae]	
713		(oxygen-evolving			cerevisiae, Peptide, 377 aa]	
	A13620	enhancer protein)	0.064	171040	[Saccharomyces cerevisiae]	0.054
		U comiono Dava c	1		· isinc)	V.U.J.
14	X06487	H.sapiens mRNA for		j	(AF020261) proline rich protein	ĺ
	7.00487	ocl2-Ig fusion gene	0.064	2429362		0.016
ı	Į	j	. [V.V.10
	h	Mus musculus cyclin-	J		l	
-	[2	lependent kinase	ł			i
			1	1		J
-)		nhibitor protein	j	k	AF082557) TRF1-interacting	
15		p15(INK4b)) gene.		a	inkyrin-related ADP-ribose	ı
	€7,5036 e	xon 2 and partial cds	0.064			le-10

CEC	Neare	st Neighbor (BlastN v	s. Genbank)	Nearest N	Neighbor (BlastX vs. Non-Redundan	Provide
SEQ					- Shoot (Blaset Vs. Non-Redundan	(Proteins)
ID	ACCESSIO	ON DESCRIPTION	PVALU	ACCERCIA		
	77	Human T cell	1	ACCESSIO	DESCRIPTION	P VAL
		receptor alpha chai	n	 		
716	U39099	mRNA, partial cds	0.063			
	1		0.005	<none></none>	<none></none>	NON
	i .	Clostridium	.1	1		
	f	acetobutylicum Kd	oC	1		
		(kdpC) gene, partia	ı	1		
		cds, sensor histidine	:	i		1
	İ	kinase homolog	ł	1	ĺ	i
- 1		(kdpD) and response	•		1	ł
- 1		regulator homolog	1	1		1
717	1100 (0-	(kdpE) genes.		1.		
/1/	U39673	complete cds Human DNA	0.063	<none></none>	<none></none>	
1					CHOILES	<none< td=""></none<>
- 1		sequence from clone 140L1 on	i	l		1
		chromosome 22q13.	.	ł		ĺ
		13.31, complete	1-	1		İ
- 1		sequence [Homo		i	(U95973) Serine	
718	AL022317	sapiens]	0.050		carboxypeptidase isolog	İ
		Japiensj	0.063	1931640	[Arabidopsis thaliana]	5.2
		Spiroplasma citri orfa				1
		and orff genes, partial			1	1
		cds, orfb, orfc, and	1		1	1
		orfe genes and	1 1		1	
- 1		Spiroplasma virus	1 1		· ·	
		SpV1-derived ORF1	! !			1
		and ORF3 genes,	1			!
		complete cds, and	1		(AF070704) envelope	İ
		SpVI-derived ORF14			glycoprotein [Human	
19	U28972	gene, partial cds.	0.063	4091939	immunodoficion and	
ł		Mus musculus limk			immunodeficiency virus type 1] (AC004877) sco-spondin-mucin-	5.2
20	HISTO	kinase (limk) mRNA,	1		like; similar to P98167 uncertain	
· <u> </u>	U15159	complete cds	0.063	3638957	[Homo sapiens]	5 !
		Homo sapiens				5.1
-	1	ipoprotein receptor-	į		1	
		elated protein	ı			
i A		LRP1), exons 39, 40, and 41	205		(AE000276) orf, hypothetical	
 		lasmodium	0.063	1788123	protein [Escherichia coli]	4.0
		alciparum	j			,
ı		hromosome 2,	[1	
1	se	ection 67 of 73 of	1]	
		e complete	ſ		1	1
2 A1		quence	0.063	22445 - 2	1_	}
			0.003	2244849	(Z97337) hypothetical protein	4.0

	\vdash	Nea	rest Neigh	ibor (BlastN vs	. Genbank)	Nearest N	Veigh	one (PleasV are N D		
		EQ	İ			335	·	oor (BlastX vs. Non-Redundant	Proteins)	
		D ACCESS		ESCRIPTION	P VALUI	E ACCESSIO	זאר	DESCRIPTION		
	-			otococcus		1.0025510	J14	DESCRIPTION (Z70203) cDNA EST	P VAL	UE
		ı		moniae methyl				EMBL D72220		
	1	- I		ferase gene	İ	ı		EMBL:D72339 comes from to gene; cDNA EST	his	
	72	7 2022	- 1	er, complete	1			EMBL:D75197 comes from the		
	-	23 L2932	3 seque	ence	0.063	3874022		gene (Caenorhahdisis alassa-1		
	1	ł	i					(2/3105) predicted using	2.3	
		ł	H.san	iens mRNA	İ	1		Genefinder: cDNA EST		i
	1	i		ling Rev-	ŀ	1		EMBL:T01277 comes from th	is	
	1			alpha > ::	1	1		gene; cDNA EST		
		1		(72632 HSREV	,	ı	ı	EMBL:T01796 comes from the	is	
		1	ERB2	H.sapiens		1 .		gene; cDNA EST		
	1	ł	mRNA	A encoding Rev	,		l l	EMBL:D32545 comes from the	is	ı
	72			lpha (internal		1		gene; cDNA EST EMBL:D33060 comes from thi	. [1
	724	X72631	fragme	ent)	0.063	3979878		gene; cDNA EST EMBL:D	l l	ı
	l	1	1,,,,				- 1	AF025467) contains similarity	1.7	4
		1.		initiation	ľ	ł	te	o drosophila DNA-binding		1
	725	U17969	comple	eIF-5A gene,			P	rotein K10 (NID:g8148)	1	1
		0.7707	Archae	oglobus	0.063	2429509	[0	Caenorhabditis elegans?	1.4	1
-		i		s section 107	1				1	1
1			of 172	of the		•			1	1
ļ	726	AE001000		te genome	0.063	3462802	(/	AF082486) nef protein [Humar	1	ı
1		1	svp[40]	=svp-related		3402802	- lin	nmunodeficiency virus type 1]	0.35	1
1		l	nuclear		Ī		- 1			1
1		1		r/retinoid	j				1	ł
			Signalin	g modulator	1		(U	J58734) weak similarity to	1	Ł
L	727	S80986	3876 nt	shes, mRNA,			an	kyrin G [Caenorhabditis		1
Γ		330500	3670 III]	<u> </u>	0.063	1326288	ele	gans]	0.093	L
ı			1		- 1		pro	oline-rich proteoglycan 2	5.055	1
			1	1	- 1		pre	ecursor, parotid - rat		ſ
			Homo sa	piens 7-60			>g	i 310200 (L17318) proline-		l
Ľ	728	AF109134		complete cds	0.063	. 1083764	FIC	h proteoglycan [Rattus		l
	- 1					1003704	nor	vegicus]	0.001	ĺ
	- 1			nRNA for	1		(AI	L021816) SPBC24E9.03c,	į	
-	729	D87466	KIAA02				unk	snown, len:251aa		ĺ
		D67466	partial cd	s	0.063	2879865	[Sc	hizosaccharomyces pombe]	6e-05	
	.		Homo sar	oiens mRNA	1		T		06-03	İ
	- 1		for KIAA		1					
7	30	AB018269		omplete cds	0.063	200525-	(AF	053455) tetraspan TM4SF		
	П				5.005	2995865	[Ho	mo sapiens]	2e-16	
	.			I	1		DDC	POTHETICAL 47.6 KD		
	ı		Cricerulus		1			OTEIN C16C10.5 IN ROMOSOME III	i	
	J		mRNA for		1			5874383 gn1 P1D e1344077	į	
7:	31		Cytochron		1		type	(RING finger)		
	<u></u> L	D86954	2А14. соп	iplete cds	0.063	2496896		norhabditis elegans)	10.22	
							•	0.024113	le-22	

-	Ne Ne	arest l	Neighbor (BlastN vs	. Genbank)		Nearest 1	Neigh	hbor (BlastX vs. Non-Redundan	
SE	- 1			P VAL	UE	ACCESSI		1	t Proteins)
<u> </u>			Plasmodium		-	ACCESSI	ON	DESCRIPTION	P VAL
1			falciparum DNA **	*					
l		- 1	SEQUENCING IN	1	- 1				
	1		PROGRESS ***		- 1			1	
732	AFOLO		from contig 4-58,	1	- 1				1
1 1 2 2	AL0102		complete sequence Mycoplasma	0.062	L	<none></none>		<none></none>	
	1		gallisepticum					ZIGITES	<non< td=""></non<>
			naemagglutinin	1	- 1			1	
			precursor genes,	1	- 1			1	
733	U9071		omplete cds	0.062	- 1	.NO.		1	
		F	Iomo sapiens clone	0.002	+-	<none></none>		<none></none>	<none< td=""></none<>
	Į	P	CL4 DNA-binding	i	1	•			1.
	ł		rotein SOX21	٠.	- 1				j
734	AFIORO		SOX21) gene,		ı		j		ľ
134	AF10704		omplete cds	0.062	_	<none></none>		<none></none>	
			aenorhabditis egans Ro					SAONES	<none:< td=""></none:<>
			Oonucleoprotein	1	1		- 1		
			toantigen mRNA,		1		- 1		
735	L41729		mplete cds	0.000	1		k	(AE000687) putative protein	
		7	piete tus	0.062		2983060	11	Aquifex aeolicus	8.6
					1			PUTATIVE	
- 1					1		100	SERINE/THREONINE-	
- 1			enorhabditis		1		l f	PROTEIN KINASE D1044.3 N CHROMOSOME III	1
- 1			gans cosmid		J			gi 495684 (U00065) contains	l
			A9D, complete		1		E	GF-like repeats; highly similar	ł
			uence enorhabditis		I		to	2C84.1; 3' exons similar to	1
36	Z99287	,	gans]	• • • •	ı		pı	rotein kinase [Caenorhabditis	ł
		Cic	34115]	0.062	<u> </u>	1176542	el	egans]	5.8
- 1		Hor	no sapiens mRNA		1				3.8
- [for 1	KIAA0614		I				
37	AB014514	prot	ein, partial cds	0.062 ⁻	,	1033395	DI	NA GYRASE SUBUNIT B	
- [Hun	nan germline			*U33393	su	bunit [Myxococcus xanthus]	3.9
- [imm	unoglobulin light	- 1			-1		_
		Chair	variable region				-		1
- 1		1.	bda-IIIb	ł			1		j
8	L29165		roup) from IgM matoid factor.				(Y	12014) RAD23 protein,	i
\top		Schis	tosoma	0.062	19	914685	isol	form II	1.3
1	[icum Chinese	ł					
-		clone						1	j
1		paran	yosin mRNA.	1					1
<u> </u>	U09364	partia	l cds.	0.062	12	350800	MI	TOCHONDRIAL	1
			um aestivum			20000	KIB	OSOMAL PROTEIN S5	1.3
			A for beta-	1			nype	othetical protein 1246 (uvrA	
	Y16242	amyla	se	0.062	7	9834	(fear	on) - Micrococcus luteus	
					20		1/11mg	anent)	0.59

	• Neare	st Neighbor (BlastN vs.	Genbank)	Negeon No.		
SE		1	Genbank,	Mediest Meig	hbor (BlastX vs. Non-Redundant F	roteins)
II	~ 1	DN DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU
<u> </u>		Leishmania pifanoi			TROPOMYOSIN I (TMI)	1. 174201
ſ		cysteine proteinase			(POLYPEPTIDE 49)	
		(cys2) gene, complet	e]	>gi 320989 pir A60607	1
74	1 M97695	cds.	0.062	1174754		0.018
1					tropomyosin - fluke (US8755) coded for by C.	0.018
- 1	1		1		elegans cDNA yk34b1.5; coded	i
ł	j		Ť	1	for by C. elegans cDNA	1
	J		1	Į.	yk13h10.5; coded for by C.	· ·
- [1		1	Į.	elegans cDNA yk46e8.5; coded	1
	I	Markenses	ł		for by C. elegans cDNA	l
	1	Methanococcus jannaschii section 68	1	·	yk46d5.5; coded for by C.	
1 _		of 150 of the		• •	elegans cDNA yk43c2.5; coded	ı
742	2 U67526		0.030		for by C. elegans cDNA	
	007520	Caenorhabditis	0.062	1330345	yk46e8	le-40
1		elegans cosmid				
	1	W09D12, complete	i i			i
		sequence			1	
ļ		[Caenorhabditis	ļ		1	ĺ
743	Z78414	elegans]	0.061	<none></none>	<none></none>	
1		Mus musculus gene			KNONES	<none></none>
1	ł	encoding filensin,			(AE000651) H. pylori predicted	
744	Y13606	exons 6, 7	0.061	2314715	coding region HP1527	4.9
}	J	1			HYPOTHETICAL 35.5 KD	
]	j				PROTEIN IN TRANSPOSON	
	Ĭ		1		TN4556 >gi 80759 pir JQ0431	
1	1	Eggplant mosaic	j		hypothetical 35.5K protein -	
745	J04374	virus genome.	0.061	141440	Streptomyces fradiae transposon	
		Marine obligately	0.081	141449	Tn4556	3.8
1	1	oligotrophic			í l	İ
	ļ	bacterium POO-10	Į		1	
	I	DNA for 16S.	1			i
		ribosomal RNA,	Į.	•	(AB000307) transcarboxylase-	ŀ
746	AB022200	partial sequence	0.061	3983593	beta	2.2
		Rat mRNA for zinc				
747	V6.000	finger protein AT-	1		(L46815) DNA binding protein	ļ
747	X54250	BP2, partial cds	0.061	1377886	Rc [Mus musculus]	0.98
		M.musculus mRNA	- 1			
748	X69942	of enhancer-trap- locus 1			(AE000748) putative protein	
740	X09942	Mus musculus mRNA	0.061	2983969	[Aquifex aeolicus]	0.57
		for scrapie responsive	- 1			
749		protein l	0.061		(AC005223) 45643	1
		H.sapiens mRNA for	0.061	4204265	[Arabidopsis thaliana]	5e-31
750		CD88 protein	0.060	<none></none>	NONT	
			0.000	MONES	<none></none>	<none></none>

	<u> </u>	Nea	rest]	Neighbor (BlastN	vs C	ienhank)		1		→ ·		
ſ,	SEQ			- Control Mastry	· s. C	Jenoank)		Nearest N	ieigh	bor (BlastX vs. Non-Redundar	nt Proteins)	_
	ID	ACCESS	ION	DESCRIPTIO	N	P VAL	UE	ACCESSIO		DESCRIPTION	·	_
_ -				7.7						T TON	P VAL	<u>.U</u>
- 1	751	U7926		Human clone 237								
	~	07920	_	mRNA, complete Plasmodium	cds	0.060)	<none></none>		<none></none>		
1	- 1			falciparum 11-1 ge						Q.O.N.E.	<non< td=""><td><u>=></u></td></non<>	<u>=></u>
	52	X0745		part I	ne							
			_	Rattus norvegicus	-	0.060		<none></none>		<none></none>	<non< td=""><td>E.</td></non<>	E.
	- 1			protein tyrosine			- 1				1 3.10.11	
	- 1			phosphatase delta	- 1		j			ĺ	j	
1 ~			18	gene, catalytic	- 1		- 1			(4.50.440.44		
/	53	U57502		lomain, partial cds	.	0.060	- 1	3452285		(AF044915) polar tube protein	n	
- 1	- [N	A:fascicularis gene	: T		7			PTP55 precursor	0.28	
75	:a	X68359		or apolipoprotein (-		ı		- 1	SHUTTLE CRAFT PROTEIN		
 '	~+	A00339		II seudomonas braB	\perp	0.060	\perp	730843	- 1	>gi 487400	1	
	- [ene for branched						84.07.400	2e-04	4
	1			hain amino acid			ı		- 1			-
				ansport carrier (LI	υL		- [- 1			1
75	5	X51634	11		"]	0.059			- 10	(U85718) CCML [Pseudomona	ıs	- 1
					┰	0.039	╁	1835622	F	outida GB-1]	8.1	1
- 1	1						1					1
- 1	-		G	ossypium hirsutum	1				- 1		1	1
	1		co	tton fiber expresse	d				- 1		1	ı
756		NF072405		otein 2 (CFE2)	1		i			lkaline phosphatase, 145K -	j	ı
1.50	+-	11-072405	m	NA, complete cds	1	0.059	L	423766	s	ynechococcus sp.	1	
	1		Sai	mbucus nigra					\top	· ·	4.7	1
	1		rib	osome inactivating					-		i	ı
1	1		pro	tein precursor	1				- [1	ı
757	A	F012899	mR	NA. complete cds		0.056	l	2442404	(A	AF034859) juvenile hormone	1	
1	1		Rat	tus norvegicus	_	0.030	-	2662481	re	sistance protein	3.3	ı
	١		hon	ner-1c mRNA,			l					ı
758	A	F093268		plete cds	(0.054		547847	1, 5	CTIN DDFG T		
1			Hyd	lra N-COL 2				347647	100	CTIN PRECURSOR	7.0	
759	١,	61046		NA for mini-								
		01040	Aral	ngen, partial cds pidopsis thaliana		0.053		<none></none>	1	<none></none>		ĺ
[]			mRN	NA for		f				4.01.12	<none></none>	
						ı			1	1	ł	
760	_AJ	005813	enzy	anthin cleavage	^	052					1	
						.052		<none></none>	<u> </u>	<none></none>	<none></none>	
		}	{ranc	dom amplified		1						
ı			hybri	dization .		- 1			JUA.	METOGENESIS	i	
l				satellite		- 1			LX.	PRESSED PROTEIN GEG	1	
- 1		ļ.	RAH	M } {Beta		J			134	>gi 2137331 pir 148361	1	
761	6-7	0047	vulga ~	ris=sugar beets.		I			Soil	GEG-154 protein - mouse 550123 (X71642)		
,01	3/	9843	Jenoi	mic, 537 nt]	0.6	025		1730145	pid o	2550123 [Mus musculus]	1	
				•				12		(ivids indscuids)	2e-16	

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-	Near	est Neighbor (BlastN v	s. Genbank)	Nearest Nai	hhar /DI . Y	
SEQ	ł			incarest ivers	hbor (BlastX vs. Non-Redundant	Proteins)
ID	ACCESSI	ON DESCRIPTION	P VALUE	ACCESSION	DE005	
		Mouse mRNA for	+	ACCESSION	DESCRIPTION	P VAL
i		GATA-2 protein,		 		
762	AB00009	6 complete cds	0.023	<none></none>		
1 1				CITONES	<none></none>	<none< td=""></none<>
		H.sapiens CpG DN.	Α.	ľ	ZINC FINGER PROTEIN 142	
763	700044	clone 67h7, forward	1	!	(KIAA0236) to Human zinc	1
703	Z62366	read cpg67h7.ftla.	0.023	3123312	finger protein(ZNF142) [Homo sapiens]	
					3-51013]	5.9
1		transmembrane glycoprotein (CD53)			hypothetical 67K protein -	
		gene, exons 2 throug	?.		Mycobacterium fortuitum	1
764	L11670	8.			plasmid pAL5000 >gi 149986	j
			0.023	<u>· 80636</u>	(M60875) ORF2	3.4
j		Sulculus diversicolor				
		DNA for IDO-like] [İ
- 1		myoglobin, complete	1 1		(AF061267) inner membrane	
765	D83984	cds	0.023	3114665	component HtxE [Pseudomonas	
- 1		S.tuberosum mRNA	3023	3114003	stutzeri]	3.4
		for inorganic	1 1			
766	Vonce	phosphate	1 1		(X02155) thyroglobulin [Bos	
/00	X98890	transporter, StPT1	0.023	683532	(taurus)	
1		Dissostichus mawsoni				1.1
- 1		preprotrypsin gene,	1			
67	U58835	complete cds	0000			
		Glomus versiforme	0.022	<none></none>	<none></none>	<none></none>
		chitin synthase gene				
68 /	AJ009630	(clone Gvchs3)	0.022	NONE	ł	
1			0.022	<none></none>	<none></none>	<none></none>
		Human gluçagon	1	ŀ		
69	J04040	mRNA, complete cds.	0.022	<none></none>	NONE	1
70		L.esculentum Asr3			<none></none>	NONE>
-	X74908	gene	0.022	<none></none>	<none></none>	
1	l.	Shigella dysenteriae	T		CHOINES	NONE>
		O-antigen]			
		polysaccharide	ł			
		piosynthesis rfbX, O-	j	1	1	j
	a	intigen polymerase]	ł	1	j
		rfc), rhamnosyl	ſ]	1	j
		ranferase I and II	1		l.	
		rfbR and rfbQ) and	l	1	ľ	1
	ri	bD genes, complete	ſ	1		- 1
L		ds.	0.022	<none></none>		1
				TIONES	<none> <</none>	IONE>

		Nea	rest	Neighbor (BlastN v	s. Genbank)	Nearest No	eighbor (BlastX vs. Non-Redundar	t Dears'	_
	SEQ ID	ACCESS			L P VAI	LUE	ACCESSIO	1		-
⊢				Mus musculus			1	DESCRIPTION	P VALI	Л
				inositol						
				polyphosphate 5-			l	1	i	
- 1	J			phosphatase II	1			I	1	
-	772	4 50 400		(INPP5P) mRNA,	1			1	1	
\vdash	12	AF04009		complete cds	0.02	2	<none></none>	<none></none>	1	
١,	73	Wasan.		H.sapiens HLA-DM	1B			THORES	<none< td=""><td>2</td></none<>	2
_ _′	''	X76776		gene	0.02	2	<none></none>	<none></none>	ALCON TO	
- [- 1			Helicobacter pylori.				- CHOINE	<none:< td=""><td>2</td></none:<>	2
	_ [strain J99 section 82	2 }			1		
1 7	74	AE00152	- 1	of 132 of the	1	- 1			ł	
<u> </u>	-	AE00132	1 10	complete genome	0.022	2	. <none></none>	<none></none>	<none></none>	
	- 1		1	A.longa rbcL, rp15.	1	j			CHOINES	4
-	- 1		In	ps8, rpl36, rps14,	1 .					
	ı			ps2, trnI,trnF, trnC	1	- 1		į.	1 .	ı
1			la	nd rpoB (partial)	1	ı		1	i	ı
1				enes > ::						ı
1	- 1			mb X75651 ALRIB	ы				}	ı
1				longa plastid genes		- 1	-		1	ı
	- 1		fo	or ribosomal	1	ı			ļ	l
1	- 1			oteins, tRNAs,	ŀ			1	1	l
1				NA polymerase		- 1		ĺ	1	ı
1	-			bunit beta and	1			}	ı,	I
77:	5	X16004		bisco large subunit	0.022		-2/04/2			ı
			La	ctococcus lactis	0.022	-	<none></none>	<none></none>	<none></none>	ı
Ī	1			emoris plasmid	ĺ					ı
1	1		pН	TW393 DNA,	I			1		ĺ
776	4_	Y12707		idii, mlladii genes	0.022	- 1	-MONT.			
1	1		Ага	abidopsis thaliana	0.022	+	<none></none>	<none></none>	<none></none>	
				tamyi-tRNA		ł		1		
777	1_	U27118	red	uctase	0.022		<none></none>]]	
1	1					1	THOMES	<none></none>	<none></none>	
	1		H.s	apiens telomeric		1		. •	1 1	
	1		DN	A sequence, clone		1		· ·	1 1	
770	١.	****		EL002, read		ł		(J05503) carbamoyl-phosphate	i 1	
778		296622	5PT	ELOO002.seq	0.022		191333	synthetase (E.C.6.3.5.5)		
	ı		١					symmetase (E.C.b.3.3.3)	9.8	
				ulus diversicolor		1				
				A for IDO-like					1	
779	_	102004		globin, complete		1		probable membrane protein		
117	۳-	83984	cds		0.022	L	1078509	YDR018c - yeast	0.7	
				picns flow-sorted			^		9.7	
				mosome 6		1	j	(AB022786) N-acetyl-beta-D-	- 1	
780	7			III fragment,		i		glucosaminidase (Enterobacter	ı	
. 00		11932	SCO	DA4A3	0.022	<u> </u>		sp.]	7.5	
					,				1.5	

-	Near	est Neighbor (Blas	N vs. C	ienbank)	Nearest	Neigh	hor (Black) we Non Baden 1	. D
SE	२					- · UIRIII	bor (BlastX vs. Non-Redundan	Proteins)
ID	ACCESS	ON DESCRIPT	'ION	P VALU	IF ACCESS	70		
		Xenopus laevi:		1 17120	E ACCESS	ION	DESCRIPTION	P VAI
		mitochondrial		 -				
781	M1021	7 complete geno		0.022	21.45	_	B2168_C2_205 protein -	
				0.022	214576	3	Mycobacterium leprae	7.3
	1	1	- 1		1		PROBABLE HELICASE	
	1		j				MOTI MOLL	1
	j		j		1	1	MOT1 Mot1p is a probable helicase essential for vegetative	
	1.	Pea chloroplast					growth on rich glucose medium	/c
	1	glyceraldehyde-	-3-				at 30 degree C. S	m
	1	phosphate	- 1			- 1	at 30 degree C: Swiss-Prot	
	İ	dehydrogenase	- 1		1	- 1	Accession number P32333; similar to S. cerevisiae RAD26	.
700		(Gpb1) gene,	j			1	gana product. Sur B	5 .
782	M55147	complete cds.		0.022	417308	ľ	gene product: Swiss-Prot	
		Acholeplasma v	irus	•	11,508		Accession number P40352 (AB008757) subunit II of	4.2
	ĺ	MV-LI DNA fo			I		c(o/b)3-type cytochrome c	1
783	V6000-	complete circula	r		1		oxidase (Bacillus	1
/03	X58839	genome		0.022	3273189		tearothermophilus)	1 .
		1				- 	VITELLOGENIN I	4.1
			- 1		1		PRECURSOR (YOLK	1
- 1			- 1		•	P	ROTEIN I)	1
- 1					[gi 72270 pir VJFF1	1
- 1		Mouse c-myb	ı		1	vi	itellogenin I precursor	1
784	M26185	oncogene, exon I	and			uı	nnamed protein product	1
	14120165	exon 2 (partial).		0.022	138592	(I	Orosophila melanogaster]	2.5
- 1		Streptomyces albi						2.3
- 1		valine dehydroger	15			- 1		1
ı		(Vdh) gene, comp	late			(A	F003145) B0414.8 gene	
85	AF061195	cds	icie	0.022		pr	oduct [Caenorhabditis	
\neg				0.022	2088768	ele	egans]	0.86
		Homo sapiens alpl	na I				ARLY GROWTH	
- 1		1,2-mannosidase I	в			RE	SPONSE PROTEIN 1 fish	
86	AF053622	gene, exon 9		0.022	1252261	>g	i 531456 (U12895) egr1	l
		S.cerevisiae	_	0.022	1352361	[[D;	anio rerio] rerio]	0.36
ı		chromosome XIV	- 1	- 1		lnı.	TATILET OF BEEN	
_	_	reading frame ORF	:	1			TATIVE TUMOR	
7	Z71500	YNL224c	- 1	0.022	1708875		PPRESSOR LUCA15	
ı					1,000/3	sap	iens]	0.16
-	ļ	Herpes simplex viri	ıs	ŀ			ĺ	
		type 2 genomic DN	A				i	
٠		for 0.74-0.84 regior	1,	- 1		(AR	0114S6) short ORF [TT	
8		omplete cds		0.022	3132276	viru		
		Zea mays T				17110	31	0.13
		cytoplasm male	- 1	- 1			Į.	1
1	S	terility restorer		- 1		(AI	031035) putative aldehyde	ŀ
	U43082 6	actor 2 (rf2) mRNA omplete cds				dehv	drogenase (Streptomyces	ľ
)				.022				

	EQ D ACCESS		Neighbor (BlastN vs.			Clenbor (BlastX vs. Non-Redundon	
F	D ACCESS			1		leighbor (BlastX vs. Non-Redundan	t Proteins)
79				P VALUE	ACCESSIO	DESCRIPTION	DVAC
79			H sapiens simple				PVALI
73	20 200		tandem repeat DNA				
	90 X8691	_	(clone wg3a6)	0.021	<none></none>	<none></none>	
1			Mus musculus	1		SNORES	<none< td=""></none<>
79) A E 100¢		Pontin52 mRNA.	1	j		1
-	1 AF1006	94	complete cds	0.021	<none></none>	<none></none>	
1	· f].	Nannostomus sp.				<none< td=""></none<>
1	1		arge subunit rRNA	The state of the s	1		
	ı		gene, mitochondrial			1	1
1	1		gene encoding	1	ĺ		- [
			nitochondrial rRNA,	ł	1		1
792	2 U34016		partial sequence.	1	,.		1.
			east mitochondrial	0.021	<none></none>	<none></none>	<none></none>
			enes for 15S rRNA	1	- 50		
793	X00845		nd tRNA-Trp	0.021	-NO.		
ĺ			lomo sapiens gene	0.021	<none></none>	<none></none>	<none></none>
	1		or CC chemokine				
	1	P.	ARC precursor.				ł
794	AB01211	3 cc	omplete cds	0.021	<none></none>		
			aucus carota		THOMES	<none></none>	<none></none>
	ł	gI	obulin-like protein	1		1	1
795			ica8) gene,	ł			1
193	U62395	co	mplete cds	0.021	<none></none>	<none></none>	1
	1					(AF004835) tyrocidine	<none></none>
796	M22710	P.1	falciparum actin II			synthetase 3 [Brevibacillus	
-750	M22718	ge	ne, complete cds.	0.021	2623773	brevis]	
		Ar	abidopsis thaliana	- 1		(AJ006631) cysteine-rich	8.8
797	U27118		tamyl-tRNA uctase			secretory protein-1 [Equus	1 1
	32.110		apiens CLN3	0.021	3549885	caballus]	8.8
798	X99832		ie, complete CDS			(\$52010) orf1 5' of EpoR [mice,	- 0.0
		1500	ic, complete CDS	0.021	262249	Peptide, 85 aa] [Mus sp.]	8.7
J		Ho	no sapiens TRAIL	- 1		SUCCINYL-	
- 1		rece	eptor 2 mRNA.	1		COA.COENZYME A	
799	AF016266		plete cds	0.021	7200	TRANSFERASE transferase	I
			-	0.021	729048	[Clostridium kluyveri]	8.7
1		1		I		LIPOPOLYSACCHARIDE 1,2-	
J			ian DNA	i			j
ł		sequ	ence from PAC	- 1		ACETYLGLUCOSAMINETR	
- 1		1791	15, BRCA2 gene	J		ANSFERASE >gi 466761]
ı	1	regio	on chromosome	1		(U00039) rfaK (Escherichia	
			2-13 contains	ł		coli] >gi 1790053 (AE000440)	J
<u>, </u>	500		se-phlorizin	i		probably hexose transferase;	1
00	Z92541	hydro	olase (LCT)	0.021	585820	lipopolysaccharide core biosynthesis	5.3

CO	Near	rest Neighbor (BlastN	I vs. Ge	enbank)	Nearest No	eighbor (BlastX vs. Non-Redundant P	
SEC	~		ON	P VALUE	1		
-		dopamine D2			1	DESCRIPTION	P VAL
	I	receptor [human				(VOSCO) NA DADA	
801	05000	brain, Genomic,			1	(Y08029) NAD(P)(+)arginine ADP-ribosyltransferase	
801	S58588		5]	0.021	2677620	[Oryctolagus cuniculus]	
	ł	Rat nerve growth	'			toryctolagus Cuniculus)	5.1
	1	factor-inducible	- 1		1		ł
802	M60522	protein (VGF) ge	ne,		j	(AF030050) replication factor C	
	14100322	complete cds.		0.021	4103934	[Rattus norvegicus]	2.1
	1	Gallus gallus	- 1				3.1
	1	neuregulin beta-1:	.		ł		
803	AF045654	mRNA, complete	oda	0.001		(AF040647) No definition line	
		inadva, complete	cus	0.021	2746829	found [Caenorhabditis elegans]	3.0
	i	1	- 1	٠. ا		(Z82056) T26H5.8	
		1	- 1			[Caenorhabditis elegans]	
804	M69023	Human globin gen		0.021	2000000	>gi 3880787 gn1 PID e1350288	
				0.021	3880259	(AL032620) T26H5.8	2.4
- 1		H.sapiens CpG DN	IA,	ł		(11000.45)	
000		clone 69d2, reverse	e			(U80845) similar to family 1 of	
805	Z65960	read cpg69d2.rt1b	.	0.021	1707245	G-protein coupled receptors	
- 1					1707243	[Caenorhabditis elegans] CORE ANTIGEN	0.79
ł		1.	- 1	- 1		>gi 73601 pir NKVLC2 core	
806	Vozoza	A.oligospora gene	- 1	1		antigen - woodchuck hepatitis	
-	X97073	encoding lectin		0.021	116949	virus 2 >gi/336135	0.47
		D. melanogaster	- 1			300 0 7 EN 300 133	0.47
		mRNA for gene containing opa	ł	}			
07	X56491	repetitive element	-1 .			HOMEOBOX PROTEIN DLX-	
		Homo sapiens		0.021	2842750	7 >gi 1620520	0.16
- [(subclone 1_f6 from	- 1				0.10
i		PI H31) DNA	1	1			
80	L78760	sequence		0.021	11267	!!!! ALU CLASS F WARNING	i
. [,.021	113671	ENTRY !!!!	0.15
	i		1	1		CYSTEINE SYNTHASE A (O-ACETYLSERINE	
- 1			1				
ł			1			SULFHYDRYLASE A) (O- ACETYLSERINE (THIOL)-	
- 1				1		LYASE A) (CSASE A)	ŀ
	1		1	1		>gi 68323 pir SYEBAC cysteine	J
- 1	j		1	1		synthase (EC 4.2.99.8) A	ı
			1	i		Salmonella typhimurium	ľ
].	Homo sapiens	j	- 1	ĺ	>gi 153935 (M21450) cysK	- 1
	B007864	KIAA0404 mRNA,		1		protein [Salmonella	[
- ^		partial cds	0.0	021		A1	1
1		Mycobacterium				y partially	0.12
1		uberculosis H37Rv	1	1	İ		
A		omplete genome; egment 22/162			ł	(AL021932) hypothetical	- 1
		-Emelit 42/102	0.0	ומ ו	2909514	protein Rv0439c	

	Neares	Neighbor (BlastN vs. (Genbank)	Nearest Nein	hbor (BlastX vs. Non-Redundant P	rotoin=1
SEC			T	- Car C31 14C12	BIASEA VS. NON-REGUNDANT P	ioteins)
В		N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-	 	Hypocrea jecorina	 			
}		mannose-1-phosphate				
		guanylyltransferase	1	ļ	(AL031538) mannose-1-	
	1	(MPGI) mRNA,	1		phosphate guanyltransferase	
811	U89991	complete cds	0.021	3581924	[Schizosaccharomyces pombe]	6e-20
		Sugar beet			i i i i i i i i i i i i i i i i i i i	00-20
1		mitochondrial				
812	X00641	minicircle pO	2 272			
F 312	700041	sequence D.melanogaster	0.020	<none></none>	<none></none>	<none></none>
1		mRNA for hdc]			
813	Z50097	protein.	0.020	<none></none>	<none></none>	-NIONTE-
1			•		THOMES	<none></none>
	J	Phoebis sennae large				
	1	subunit ribosomal RNA gene, partial	ļ l			
ł	1	sequence; tRNA-Val				
		gene, complete			1	
		sequence; and small				
1	'	subunit ribosomal				
]	RNA gene, partial	l l			
		sequence,				
	f	mitochondrial genes	1		ŀ	
814	A FO 44066	for mitochondrial	[•	l	
814	AF044866	RNAs	0.020	<none></none>	<none></none>	<none></none>
		Sambucus nigra	ſ		l i	
		hevein-like protein				1
815	AF074386	mRNA, complete cds	0.020	<none></none>	<none></none>	<none></none>
					4.01.2	714014E2
		Arabidopsis thaliana				
	.	cellulose synthase	-8-			. !
		catalytic subunit (Ath-B) mRNA, complete		•]	1
816		cds cds	0.020	<none></none>	1	
		Plasmodium	0.020	CHOINES	<none></none>	<none></none>
		falciparum	j		j i	ł
ľ		chromosome 2,	j			1
1		section 42 of 73 of	1		(AF003342) bunched gene	Ì
817		the complete			product [Drosophila	
01/	AE001405	sequence	0.020	2196776	melanogaster]	8.4
İ		Sambucus nigra	1			
i		nevein-like protein	1		histidine-rich protein -	
818		nRNA, complete cds	0.020	627071	Plasmodium lophurae	2.8
						2.0

	Nearest	Neighbor (BlastN vs. C	ionhank)	None		
950		THEIRIOU (BIASUA VS. C	enoank)	Nearest Neig	hbor (BlastX vs. Non-Redundant P	rotcins)
SEQ	· I		•	1		
	ACCESSION	1 DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE
1		Hylobates hoolock				1
1		mitochondrial DNA	ł	i	(DI0043) ORF [Acetobacter	
819	Y13304	for cyth gene, Horace	0.020	285580	pasteurianus)	1.2
1	Í	<u> </u>			(U80439) coded for by C.	
ļ					elegans cDNA yk7c8.5; coded	ļ
	<u> </u>	1			for by C. elegans cDNA	i
1	ł				yk I33b3.5; coded for by C.	l i
1					elegans cDNA yk65a4.5; coded	,
Į.	İ	1			for by C. elegans cDNA]
l I		1			yk7c8.3; coded for by C.	
	- 1	77			elegans cDNA CEESQ66F;	:
820	Z 66539	H.sapiens creatine			coded for by C. elegans cDNA	
620	200339	transporter gene	0.020	1703594	yk65a4.3;	0.98
1 1			i		EARLY GROWTH	
l i		Homo sapiens alpha I.2-mannosidase IB			RESPONSE PROTEIN I fish	
821	AF053622	gene, exon 9	0.000		>gi 53 I456 (U12895) egr1	
021	AI 033022	gene, exon 9	0.020	1352361	[Danio rerio] rerio] HYPOTHETICAL 38.1 KD	0.72
					PROTEIN IN SBCB-HISL	j
			I		INTERGENIC REGION	
			1		>gi 405956 (U00009)	
			i		ORF_ID:0349#4; similar to	
1			ı		[SwissProt Accession Number	ľ
		•			P33015] [Escherichia coli]	
f		Í	ľ		>gi 1736693 gnl PID d1016570	- 1
- 1			1	·	Number P33015] (Escherichia	
- 1		Human MHC class II	- 1		coli] >gi 1788323 (AE000292)	,
j		HLA-DRw53-beta	i		putative transport system	
		(DR4,w4) gene,			permease protein [Escherichia	}
822	M20555	exons 2,3,4,5,6.	0.020	465569	coli]	0.43
T		Human MHC class II			COENZYME PQQ	0.43
1	ļ:	HLA-DRw53-beta	1		SYNTHESIS PROTEIN F	į
J		(DR4,w4) gene,	. [synthesis F - Pseudomonas	
823	M20555	exons 2,3,4,5,6.	0.020	1709751	fluorescens >gi 929802	0.42

	Ne	arest i	Veighbor (BlastN v	s. Genbank)		Nearest I	Veigh	bor (BlastX vs. Non-Redundant	Protei
SE	-	SION	DESCRIPTIO	N P VAL	UE	ACCESSI(DESCRIPTION	P VAL
	 -			_ <u> </u>				INTOTACTICAL IZED KU	TI VAL
								PROTEIN ZK637.2 IN CHROMOSOME III >gi 102507 pir S15787 hypothetical protein 1 (cosmid	
								ZK637) - Caenorhabditis elegans Genefinder; cDNA EST yk217b5.3 comes from this gene; cDNA EST yk217b5.5	
		H	Iomo sapiens mRN	IA				comes from this gene; cDNA EST yk340g12.3 comes from this gene; cDNA EST	
824	AJ0050	fo	or putative SMC-li otein, partial		1	267449	.	yk340g12.5 comes from this gene; cDNA EST yk428e5.5	le-12
825	AF03409	gi sy	occaria bicolor yoxal malate nthase protein				(U41538) No definition line	.0-12
023	Ar03409	_	RNA, complete cd	5 0.020		1109847	f	ound [Caenorhabditis elegans]	1. 22
826	AF10069	Po	ntin52 mRNA, mplete cds	0.019		132836	I.	OS RIBOSOMAL PROTEIN 28 protein L28 [Rattus	le-22
827	AF09326	hor	ttus norvegicus ner-1c mRNA, nplete cds	0.019	\dagger		(2	orvegicus] 299109) similar to DNA	5.7
28	AF100694	Mu Por	s musculus ptin52 mRNA, aplete eds		T	2633401	\top	CONUCTED SESSISTANCE	4.5
	111 100074	Con	ipiete cas	0.019	╁╴	2492604	PF PC	ROTEIN CDR2 albicans]	4.4
		1	ı				CI	LUCOSYLTRANSFERASE 7C9.07	
			·				(Z:	i 1314159 gn PID e241760 73099) SPAC17C9.07,	·
		janna	nanococcus aschii section 80 60 of the				50 SW	ative glucosyl transferase len: l, similar to V:ALG8_YEAST P40351	
9	U67538	COmp Hum	olete genome an periodic	0.019		1723566	glu	cosyltransferase ALG8	2.7
0	U56088	trypto	ophan protein 2 2) gene, exons	0.019		2144804	coll bov	agen alpha 1(II) chain -	
	U76524	ribosc protei	ucus nigra ome inactivating n precursor				(U9	1682) vitciline membrane ein homolog [Aedes	0.040
	070324	mKN/	A. complete cds	810.0		1916976	aegy		7.2

SEQ ID		Nearest	Neighbor (BlastN vs. (Jenbank)	Nearon Nai	hhor (Plant V us Now Posture	
D ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION P VALUE ACCESSION DESCRIPTION ACCESSION DESCRIPTION	SEO		Total Total		Memer Melo	moor (Blasta vs. Non-Redundant Pi	roteins)
Onobrychis vicilifolia chalcone synthase (CHS) mRNA, complete eds			DESCRIPTION	DVALIE	10000000		
Chalcone synthase (CHS) mRNA. CMS Sequence Sequence CHS MRNA. Complete cds CHS MRNA. Complete cds CHS MRNA. CMS CHS CMS		, reception	DESCRIPTION .	T VALUE	ACCESSION	DESCRIPTION	P VALU
Calcone synthase (CHS) mRNA, Complete cds CHS) mRNA, Complete cds CHS) mRNA, Complete cds CHS) mRNA, Complete cds CHS) mRNA, CHSP (XL- INCENP) mRNA, CHS	 		Onobrychie viciifolia			(748700) 770 KG	
S32 AF026258 CHS) mRNA, Complete eds O.018 763076 Carpitol S.			1	1 .	'		1
832 AF026258 complete cds Co				Ī			}
Name Name	832	AF026258	T	0010	7/207/		1
INCENP (XL- INCENP) mRNA,	002	74 020236		0.018	/630/6	carpio	5.2
S33				}			İ
833		·				(A 1006 428) have	j
C.jejuni VSI DNA >	833	U95094	I '	0.000	3055011		
Sequence 2 from Patent WO9417205 >				0.009	293011	В	0.60
Sequence 2 from Patent WO9417205 > :: gb]T6090 T6090 Sequence 2 from Patent WO9417205 > :: gb]T6090 T6090 Sequence 2 from patent US 5691138 0.008 NONE			::	!			
Sequence 2 from Patent WO9417205 >			emb[A39603]A39603	i i	1,		
Patent WO9417205			•				
Sade				٠.			
Sequence 2 from patent US 5691138							ľ
Rattus norvegicus HEAD FIBER PROTEIN (LATE PROTEIN GP8.5) Sgi[75846 pir WMBP8H gene S.5 protein - phage PZA Sgi[216057 (M11813) head fiber protein SUPEROXIDE DISMUTASE (FE) 1.15.1.1) (Fe) - Pseudomonas aeruginosa Sgi[409767 S.9			_ ,				
Rattus norvegicus Rattus norvegicus homer-1c mRNA, complete cds 0.008 138116 SUPEROXIDE DISMUTASE (FE) 1.15.1.1) (Fe) - Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1711436 Pseudomonas aeruginosa 1811436 Pseudomonas aeruginosa	834	X71603	•	0.008	<none></none>	NONE	-NIONITA
Rattus norvegicus homer-1c mRNA, complete cds			<u> </u>				CHONES
Rattus norvegicus homer-1c mRNA, complete cds							
Rattus norvegicus homer-1c mRNA, complete cds 0.008 138116	ĺ						
Nomer-1c mRNA, complete cds 0.008 138116	J		Rattus norvegicus	i			
R35			homer-1c mRNA,	<u> </u>			
Bovine herpesvirus type 1 UL7 gene	835	AF093268	complete cds	0.008	138116	fiber protein	8.1
Bovine herpesvirus type I UL7 gene 0.008 1711436 >gil409767 5.9	1					SUPEROXIDE DISMUTASE	
Sacharomyces cerevisiae Sacharomyces cer	ł						
Arabidopsis thaliana	006					Pseudomonas aeruginosa	
1-aminocyclopropane 1-carboxylate synthase (ACS2) gene, complete cds. 0.008 683698 [Saccharomyces cerevisiae] 1e-0e	830	X91751	type I UL7 gene	0.008	1711436	>gi 409767	5 .9
1-aminocyclopropane 1-carboxylate synthase (ACS2) gene, complete cds. 0.008 683698 [Saccharomyces cerevisiae] 1e-0e	ł	İ	Archidonaia shaliana	l			
1-carboxylate synthase (ACS2) gene, complete cds. 0.008 683698 [Saccharomyces cerevisiae] 1e-06	- 1			j			
Synthase (ACS2) gene, complete cds. 0.008 683698 [Saccharomyces cerevisiae] 1e-06	1			İ		1	
837 M95594 gene, complete cds. 0.008 683698 [Saccharomyces cerevisiae] 1e-06	- 1			1			
Methanococcus jannaschii section 7 of 150 of the (Z68493) predicted using 1e-00	837	- 1		0.000	. (02/00		
jannaschii section 7 of 150 of the (Z68493) predicted using (Z68493) predicted using 1e-07 (Z68493) predicted				0.008	84000	[[Saccharomyces cerevisiae]	1e-06
Sample Of 150 of the complete genome O.008 3874664 Genefinder 1e-07	- 1			1			
838 U67465 complete genome 0.008 3874664 Genefinder 1e-07 839 X72388 B.taurus mRNA for filensin 0.008 100174 Carboxylate synthase 7e-09 840 U22398 mRNA complete cds 0.008 2228750 Subunit [Homo sapiens] 2e-18 840 U22398 MRNA complete cds 0.008 2228750 Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 840 U22398 RNA polymerase III Subunit [Homo sapiens] 2e-18 3e-18	1	P		1		(768402) prodicted ::::	
B.taurus mRNA for filensin	838			0.008	3874664		1. 67
No.008 N				0.000	J01+004		1e-0/
Human Cdk-inhibitor p57KIP2 (KIP2) WRNA. complete cds. 0.008 2228750 (U93868) RNA polymerase III subunit [Homo sapiens] 2e-18	839	1		0.008	100174		72.00
P57KIP2 (KIP2) (U93868) RNA polymerase III subunit [Homo sapiens] 2e-18			·		100174	carooxylate synthase	76-09
P57KIP2 (KIP2) (U93868) RNA polymerase III subunit [Homo sapiens] 2e-18	- 1	ļ.	Juman Cdk-inhibitor	Ì			j
840 U22398 mRNA, complete cds 0.008 2228750 subunit [Homo sapiens] 2e-15	l			j		(U93868) RNA polymerase III	
Xenopus laevis LIM	840	4.5		0.008	2228750		20-15
						The same of the sa	76-19
			lass homeodomain	ļ]	
841 L42546 protein 0.007 <none> <none> <none< td=""><td>841</td><td>L42546 p</td><td>rotein</td><td>0.007</td><td><none></none></td><td><none></none></td><td><none></none></td></none<></none></none>	841	L42546 p	rotein	0.007	<none></none>	<none></none>	<none></none>

	Nearest I	Neighbor (BlastN vs. G	enbank)	Nearest Neighb	oor (BlastX vs. Non-Redundant Pr	roteins)
SEQ						
I ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Homo sapiens		11000301014		I VALUE
—		ribosomal protein s4				
Į i		X isoform gene,				
842	AF041428	complete cds	0.007	<none></none>	<none></none>	<none></none>
100	74 041420	Secale cereale omega	0.007	CHOILES	GNONES	KNONES
1		secalin gene.				
843	AF000227	complete cds	0.007	<none></none>	<none></none>	<none></none>
J.J.	111 000227	Human MHC (HLA)	0.007	CHOILE	- GROTIES	- CINOINE
		DRB intron 1 DNA,				ĺ :
844	D86254	partial sequence	0.007	<none></none>	<none></none>	<none></none>
		Sambucus nigra		• •		1
		ribosome inactivating			}	1
		protein precursor		÷]
845	AF012899	mRNA, complete cds	0.007	<none></none>	<none></none>	<none></none>
1		M.musculus gene for				
846	Y07738	vimentin	<u>0</u> .007	<none></none>	<none></none>	<none></none>
1		Arabidopsis thaliana				<u> </u>
		mRNA for				
		neoxanthin cleavage		.02		
847	AJ005813	enzyme	0.007	<none></none>	<none></none>	<none></none>
		Homo sapiens alpha-	i			
848	AFOSSIIO	tectorin (TECTA)	0.007	NONE	NONE	NONT
040	AF055119	gene, exon 6	0.007	<none></none>	<none> .</none>	<none></none>
1 1		Zucchini 1-				
1	i	aminocyclopropane-1				f i
849		carboxylate synthase	0.007	<none></none>	<none></none>	<none></none>
		Homo sapiens DSG3				
		gene, partial intron				
		and partial exon 6,	ŀ			1
850		140 bp	0.007	<none></none>	<none></none>	<none></none>
		M.voltae vhuD,				
		vhuG, vhuA, vhuU &	ľ			
851		vhuB genes	0.007	<none></none>	<none></none>	<none></none>
		Brassica rapa mRNA				
l		for SLG45, complete				
852	AB012105	cds	0.007	<none></none>	<none></none>	<none></none>
ŀ			ļ			
ĺ	l l	telomere:	i			
		(minichromosome,	J	ĺ		ļ l
İ		repeats}		ļ		[]
853		[Trypanosomá brucei.]	0.007	NONE	ANONIT	NOST
633	S43882	Genomic, 1170 nt]	0.007	<none></none>	<none></none>	<none></none>

	Nearest	Neighbor (BlastN vs. C	ienbank)	Nearest Noish	bor (BlastX vs. Non-Redundant Pr	oteins)
SEO		13.00. (3.030. 13.0		Tromest Meigh	Wilder VS. Non-Redundant Pr	O(CIUS)
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE
		 	1	1 HEEZSSION	J DESCRIPTION	I VALUE
		Geomydoecus nadleri		 		
		mitochondrial	1			l
1		cytochrome oxidase I			1	İ
854	L32674	gene, partial cds.	0.007	<none></none>	<none></none>	<none></none>
į i		Caenorhabditis				
		elegans cosmid			1	
855	U58732	F48D6.	0.007	<none></none>	<none></none>	<none></none>
ì						
		Sambucus nigra				Í
		ribosome inactivating		٠.		-
856	U76524	protein precursor mRNA, complete cds	0.007	-NONT-	NOVE	
1000	070324	marya, complete cus	0.007.	<none></none>	<none> HYPOTHETICAL 121.1 KD</none>	<none></none>
1				1	PROTEIN IN BIO3-HXT17	
		H.sapiens mRNA for			INTERGENIC REGION	
		MDR3 P-			PRECURSOR YNR067c - yeast	
857	Z35284	glycoprotein	0.007	1730696	(Saccharomyces cerevisiae)	9.5
					(Outerationity ets estevisiae)	7.5
		Human sno oncogene			<u>j</u>	
		mRNA for snoA			(U24203) membrane protein	
858	X15217	protein, ski-related	0.007	902455	[Escherichia coli]	8.8
		Arabidopsis thaliana			1	
		cellulose synthase				1
		catalytic subunit (Ath- A) mRNA, complete	٠		arous a open is	ı
859		cds	0.007	1684636	(Y09454) ORF3 (Lactobacillus	
992	74 OZ/173	cus	0.007	1004030	casei bacteriophage A2]	8.3
	l	Sambucus nigra				ŀ
ł		ribosome inactivating				
		protein precursor			(Z48795).R05H5.7	i
860	AF012899	mRNA, complete cds	0.007	3878803	[Caenorhabditis elegans]	8.3
I		119=180-200 kda				
		membrane protein				
1		scavenger receptor				
Í		homolog (clone 18,				ļ
		intron and flanking				
i		exons 14 and 15}				
l	4	sheep, lymph node,	ŀ			
İ		ymphocytes, Genomic, 308 nt.			(I 08174) ODE2	ļ
861		segment 2 of 2]	0.007	294747	(L08174) ORF2	٠, ١
	3,0317	registers = U(a)	0.007	474/4/	[Romanomermis culicivorax]	7.4

	:÷.	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant P	
	SEQ				Treatest (ver	gnoor (Blastix vs. Non-Redundant P	roteins)
L	ID	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	1
					i iteezosioit	DESCRIPTION	PVALUE
			Pedicularis				
		ļ	verticillata	ł	j		
			chloroplast DNA,	ł			İ
		j	intergenic region	1	j		Ĭ
	862	B0000.	between trnT(UGU)		Ī	(AF026789) vitellogenin	
-	802	D88084	and trnL(UAA)5'exc	п 0.007	2555187	[Pimpla nipponica]	6.9
			Chicken mRNA for	1		CD30L RECEPTOR	
			aldehyde	1		PRECURSOR	
	863	X58869	dehydrogenase	0.007		(LYMPHOCYTE	
		1150005	denydrogenase	0.007	115978	ACTIVATION ANTIGEN	. 6.5
1	I			1		cDNA EST EMBL:D35637	
				'		comes from this gene; cDNA	1 .
	- 1		1	1		EST yk322a3.5 comes from this	
	1]]		gene; cDNA EST yk397b2.5]
	- 1		1]]		comes from this gene; cDNA	
	- 1		1	1		EST yk348b11.5 comes from	
1	ı		•	1 1		this gene; cDNA EST	
	- 1		j	1 1		yk397b2.3 comes fr	
	- 1			[>gi 3880965 gn1 PID e1350578	
	- 1			i		comes from this gene; cDNA	
1	ı			1		EST yk322a3.5 comes from this	
				1 1		gene; cDNA EST yk397b2.5	ŀ
	- 1		Homo sapiens mRNA	i 1		comes from this gene; cDNA	1
			for GS3786, complete			EST yk348b11.5 comes from	
8	64	D87120	cds	0.007	3879589	this gene; cDNA EST yk397b2.3 comes	
			H.sapiens gene for	0.00	3079309	(AF010403) ALR [Homo	5.1
80	55	X68793	antithrombin III	0.007	2358285	(Au 010403) AER [Homo	20
						HYPOTHETICAL 29.8 KD	3.8
l						PROTEIN IN HOLB-PTSG	i
1	1	-		i	•	INTERGENIC REGION	1
1				1		>gi 1787342 (AE000210) orf,	· [
	ı		Danio rerio mRNA	1		hypothetical protein	
	-		for opioid receptor	1		[Escherichia coli] protein in	•
86	6		homologue	0.007	2507509	holB 3'region . [Escherichia	
_	T			0.007	230/309	[coli]	1.9
			Streptomyces albus	1		1	
	1		valine dehydrogenase	1		(AF003145) B0414.8 gene	Ì
	_ [(Vdh) gene, complete	1		product [Caenorhabditis	ſ
86	7 /	AF061195	eds	0.007	2088768	elegans]	1.9
	1		A make indicate the second second			UDP-N-	
			Arabidopsis thaliana	ļ		ACETYLGLUCOSAMINE 2-	·
	1	1	nRNA for	j		EPIMERASE UDP-N-	
868	3 4		neoxanthin cleavage	0.007	1710.	acetylglucosamine 2-epimerase	- 1
	<u> </u>		IIIC	0.007	1710105	[Plasmid pWQ799]	1.7

		37 1 1 1 1 1 1 1 1 1				
		Neighbor (BlastN vs. (Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant Pi	oteins)
SEC	`	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	
	7	Zebrafish retinoic		I I CCL STON		P VALUE
		acid receptor alpha				<u> </u>
869	L03398	2.A	0.007	2239219	(707210) humah si at	0.77
		Human mRNA for	9.007	2439219	(Z97210) hypothetical protein	0.77
		KIAA0150 gene,]	İ	(Z14014) Pistil extensin like	
870	D63484	partial cds	0.007	19917	protein, partial CDS only	0.61
871	M31483	Maize glyceraldehyde 3-phosphate dehydrogenase, 3' end.	0.007	543068	mucin, tracheobronchial - dog	0.45
872	AF090115	Lycopersicon esculentum cytosolic class II small heat shock protein HCT2 (HSP17.4) mRNA, complete cds	0.007	2494941	ALPHA-2B ADRENERGIC RECEPTOR adrenoceptor [Cavia porcellus] >gi 1587159 prf 2206293B adrenoceptor alpha2B [Cavia	
	10000113	complete eds	0.007	2494941	porcel[us]	0.42
873	AF064029	Helianthus tuberosus lectin 1 mRNA, complete cds	0.007	1110587	(S79410) nuclear localization of signals Peptide, 140 aa] [Mus sp.]	0.26
874	X88931	H.sapiens PAL2A	0.007	1706176	CUTINASE TRANSCRIPTION FACTOR 1 ALPHA >gi 1262912 (U51671) cutinase transcription factor 1 [Fusarium solani f. sp. pisi]	0.21
875		zRAR alpha =retinoic acid receptor alpha [zebrafish, embryos, mRNA, 1773 nt]	0.007	2239219	(Z97210) hypothetical protein	0.11
876	M74193	Petromyzon marinus plasma albumin mRNA, complete cds.	0.007	730888	OCTAPEPTIDE-REPEAT PROTEIN T2	0.011
877		Saccharomyces cerevisiae Spp41p (SPP41) gene. complete cds.	0.007	3820885	(AL033126) 65G3.k [Drosophila melanogaster]	0.001
878	t	Homo sapiens mRNA for Laminin-5 beta3 thain, complete cds	0.007		(X96713) collagen [Globodera pallida]	3e-06

	<u> </u>	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant I	Dentai 1
	SEQ					and (Blasta Vs. 14011-Redundant I	roteins)
	ID	ACCESSIO		P VALUE	ACCESSION	DESCRIPTION	P VALUE
			Caenorhabditis				I VALUE
ı			elegans putative		<u> </u>		
- 1		1	transcription factor	I	1		1
ı			MAB-3 (mab-3)			(AF095741) unknown [Rattus]
Ļ	879	AF022388		0.007	3747107	norvegicus]	
- [Acanthamoeba			· Horvegicus)	5e-09
- 1		ı	castellanii	1			1 1
- 1		ł	transformation-	1	1		1 1
-1		ľ	sensitive protein	1		1	1 1
-			homolog mRNA,	1 1		(U89984) transformation-	i i
L	880	U89984	complete cds	0.007	. 1890281		2-00
Т						sensitive protein homolog	2e-09
						rabGAP domains; cDNA EST	1
1						EMBL:D34945 comes from this	1
1				1		gene; cDNA EST	1 1
Ĺ	- 1		•	1 1		EMBL:D27313 comes from this]
	l			1		gene; cDNA EST	1 1
]		EMBL:D34829 comes from this	1 1
1			1	l 1		gene; cDNA EST	i i
1				1 1		EMBL:D27312 comes from this	1
1	ı]		gene; cDNA Probable	1 1
1	ł		İ			rabGAP domains; cDNA EST	
	- 1			1		EMBL:D34945 comes from this	
1	- 1		1	1		gene; cDNA EST	1
1						EMBL:D27313 comes from this	! !
	- 1		j,			gene; cDNA EST	
l						EMBL:D34829 comes from this	
			Homo sapiens mRNA			gene; cDNA EST	1
۱.	81	4 D000 c00	for KIAA0882			EMBL:D27312 comes from this	Į.
<u>ا</u>	01	AB020689	protein, partial cds	0.007	3880809	gene; cDNA	1e-23
l			Mus musculus				
۱۵	82	AF100694	Pontin52 mRNA,				
۲	"-	A1100094	complete cds	0.006	<none></none>	<none></none>	<none></none>
ĺ			Arabidopsis thaliana				
			cellulose synthase				Ī
			catalytic subunit (Ath-		·		ŀ
				l l			1
88	33		A) mRNA, complete cds	0.00-			1
	~	11102/1/3	cus	0.006	<none></none>	<none></none>	<none></none>
	1		Sambucus nigra	1			
	- 1		ribosome inactivating	1	,	ļ	ł
			protein precursor	ſ	į		ŀ
88	4	* ***	nRNA, complete cds	0.006	.,,,,,,	- 1	
		3,0024	mara, complete cus	0.006	<none></none>	<none></none>	<none></none>

	Near	est Neighbor (BlastN	vs. Genbank)	Negroe N.	inhhar (D)	
S	EQ			Nearest Ne	ighbor (BlastX vs. Non-Redundar	t Proteins)
	D ACCESSI	ON DESCRIPTION	N PVALU	E ACCESSION	N DESCRIPTION	P VALUE
 						FVALUE
- 1	ı	Sambucus nigra				
- 1	1	ribosome inactiva				İ
1.		protein precursor			}	- 1
88	35 U76524	mRNA, complete	cds 0.006	<none></none>	MONE	
		Arabidopsis thalia		1	<none></none>	<none></none>
- 1		mRNA for	1			
		neoxanthin cleava	ige		1	1 1
88	6 AJ00581	3 enzyme	0.006	<none></none>		1
- 1	1	Brassica rapa mRI		CHOILS	<none></none>	<none></none>
j		for SRK45, compl		1.		
88	7 AB01210		0.006	<none></none>		1 .
		Rattus norvegicus	0.000	KNONES	<none></none>	<none></none>
- 1		ceruloplasmin gene	e		ĺ	
888	M80529	exon I and 5' flank	0.006	NONE	1	
			0.000	<none></none>	<none></none>	<none></none>
ļ		Arabidopsis thaliar	na I		han a see a see	
i	1	cellulose synthase			hypothetical protein 6 -	
- 1	1	catalytic subunit (A	\th.		Chlamydomonas reinhardtii	1 1
1		A) mRNA, complete	te .	1	transposon	
889	AF027173	cds	0.006		>gi 1360717 gn1 PID e33461] [
			0.006	99408	reinhardtii]	9.6
1	ł	Sambucus nigra lec	tin	1		
		precursor mRNA,	```	ł		1 1
890	U76523	complete cds	0.006	400000	(AF039110) polyprotein	1 1
		Rattus norvegicus	0.000	4039024	[Rubella virus]	9.3
1	ł	homer-1c mRNA,		·		
891	AF093268	complete cds	0.006	160522	(M94428) merozoite surface	1 1
		Tours out	0.000	160533	antigen I [Plasmodium vivax]	7.5
1	í	Brassica rapa mRNA	. i			
	l	for SRK45, complete	,		(AF093984) envelope	1
892	AB012106	cds		40.5:-:	glycoprotein [Human	f
	12.50	Arabidopsis thaliana	0.006	4019458	immunodeficiency virus type 1]	7.0
1	ł	mRNA for	1 1			
1		neoxanthin cleavage	1		(U91682) vitelline membrane	1 1
893	AJ005813	enzyme			protein homolog [Aedes	1
			0.006	1916976	aegypti]	6.8
1 1			1		promastigote surface antigen-2	
		Rattus norvegicus]		(clone 4.6) - Leishmania major] <u> </u>
		homer-1c mRNA,	1		(fragment) >gi 9583 (X57135)	1
894	AF093268	complete cds	1		surface antigen P2 (Leishmania	1 1
	0/2/208	Rattus norvegicus	0.006	102059	major]	2.4
			1 1			
895	AF093268	homer-1c mRNA,			(AF067204) transcription factor	l I
	14 073208	complete cds	0.006	3171241	BF-1 [Danio rerio]	1.0
896	X99384	M.musculus mRNA				
	77730+	for paladin gene	0.003	<none></none>	· <none></none>	<none></none>

		Nea	rest Ne	ighbor (BlastN vs.	Genbank)	Non		
i	SE	2				ivearest Nei	ghbor (BlastX vs. Non-Redundant	Proteins)
	ΙĐ		ION	DESCRIPTION	P VALUE	1		P VALU
ł				enhido sie d'al				IL VALU
J		i		rabidopsis thaliana				
		1		Ilulose synthase	1			
- 1		1	ca	talytic subunit (At	h-∱	1	Ĭ	1
1	897	A 5005.		mRNA, complete	İ			1
F	097	AF0271	74 cd	<u>S</u>	0.003	<none></none>	- NONE>	2702
		1	P-				THORES	<none></none>
- 1		1		rrelia burgdorferi	1	1		1
- I	898	AE00114		ction 34 of 70) of	1		(AJ011856) ORF Q0255	1
-	070	AE00114	8 the	complete genome	0.003	4160388	[Saccharomyces cerevisiae]	7.6
		1	ا ا	alada a ar ar ar			control of the state of the sta	7.6
		l		bidopsis thaliana		'	j	
		j		lulose synthase	1	1	NUCLEAR ENVELOPE PORE	-1
- 1		1		alytic subunit (Ath			MEMBRANE PROTEIN POM	1
- 1,	899	AEOGZIZ	, JA) I	mRNA, complete]	Î	121 (PORE MEMBRANE	ł
 '	033	AF02717	3 cds		0.003	1709213	PROTEIN OF 121 KD) (P145)	1 , ,
	į		1				(F143)	1.5
- 1				opersicon			1 ,	1
- 1	- 1			ilentum class II			1	
	1			ll heat shock			1	
١	000	U72396		ein Le-HSP17.6			•	
<u> </u>		072396	mKi	NA, complete cds	0.002	<none></none>	<none></none>	<none></none>
i	I			musculus	1		4.0.12	CIAOMES
٥	01 l	_AF100694		in52 mRNA,	i		l j	
1	`` +	AF100094		plete cds	0.002	<none></none>	<none></none>	-NONE.
1				mydomonas			1	<none></none>
1	- 1			ardtii light	1			1
1	- 1		narve	sting complex II	Į.		1	1
1				in precursor	- 1			I
90	22	AF104631		o3) mRNA,	- 1			
1	-	· tr 104031		lete cds	0.002	<none></none>	<none></none>	<none></none>
1				musculus	T			710115
90	3	AF100694		n52 mRNA,	1			
	+	74°100094	comp	lete cds	0.002	<none></none>	<none></none>	<none></none>
l	- 1		for Co	ica rapa mRNA				7.1011E>
904	4	AB012106		KK45, complete	1	I		
ٽٽ ا	- -	12012100	cds	n non-histone	0.002	<none></none>	<none></none>	NONE>
1								STONES
	- 1		LIMC	osomal protein	1	1	İ	
905	5 1	M21339	LIMIG-	14 gene,		1	· 1	ı
	+	11121339	comple	ete cds.	0.002	<none></none>	<none></none>	NONE>
	1		Sumbu	gua sissa	1			TAOIATS
	1			cus nigra	ł	1		1
				ne inactivating	i		1	Ī
906				precursor	1			
		11 012099	MKINA	, complete cds	0.002	<none></none>	<none> <</none>	NONE>

CE	Nea	rest Neighbor (BlastN vs.	. Genbank)	Nearest Neig	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC	-				DESCRIPTION			
 		Human h-lys gene for	or	1	DESCRIPTION	P VAI		
907	V5710	lysozyme (upstream						
-207	X5710	3 region)	0.002	<none></none>	<none></none>	1		
	1	Sambucus nigra	1		3.000	<non< td=""></non<>		
	1	hevein-like protein	1	1	i			
908	AF07438	mRNA. complete cd	0.000	}				
		Human CD4	s 0.002	<none></none>	<none></none>	<non< td=""></non<>		
	I	promoter, partial				1.01.		
909	U01066		0.002	NO	1	İ		
		Barley mRNA	0.002	<none></none>	<none></none>	<non< td=""></non<>		
910	L28094	sequence.	0.002	ANIONIE				
	i	Homo sapiens DNA		<none></none>	<none></none>	<non< td=""></non<>		
	1	from chromosome 19	- '					
		cosmid f19399 (-17	. !			1		
۸.,		kb EcoR1 restriction	j i		Í	1		
911	AD00083	fragment)	0.002	<none></none>	<none></none>			
- 1					ANONES	<none< td=""></none<>		
- 1		Homo sapiens TRHR	i 1			1		
912	AJ011701	gene promoter and	1			1		
	73011701	exons 1-2, partial Mus musculus	0.002	<none></none>	<none></none>	NONE		
- 1		Pontin52 mRNA,				KNONE		
913	AFI00694	complete cds	0.000					
		Junipiere eus	0.002	<none></none>	<none></none>	<none< td=""></none<>		
İ		Homo sapiens retinol	- 1	İ				
- [dehydrogenase gene,	1			1		
14	AF037062	complete cds	0.002	<none></none>				
1		Rattus norvegicus		CHOIVES	<none></none>	<none:< td=""></none:<>		
. ا ء.		homer-1c mRNA,	1			1		
15	AF093268	complete cds	0.002	<none></none>	AIONE	ĺ		
		Methanococcus			<none></none>	<none></none>		
1		jannaschii section 150				1		
6	U67608	of 150 of the			·			
+	307008	complete genome	0.002	<none></none>	<none> .</none>	<none></none>		
-		Arabidopsis thaliana	1			TONES		
		cellulose synthase	1	l				
-]		catalytic subunit (Ath-	j	į	•			
-		A) mRNA, complete	j	ł				
7 A		cds	0.002	NOVE				
		H.sapiens DNA for	0.002	<none></none>	<none></none>	<none></none>		
		repeat region (ABM-	- !	1				
3	Z46736	C82)	0.002	NONE				
		Brassica rapa mRNA		<none></none>	<none></none>	<none></none>		
					·			
		or SRK45, complete	- 1	j		j 1		

	Nearest	Neighbor (BlastN vs. C	ienbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	PVALUE	
		X.laevis mRNA for		cczsololv	- DESCRIPTION	I VALUE	
920	Z85983	NOVA protein	0.002	<none></none>	<none></none>	ANONTES	
	203703	Tro vis protein	0.002	CHOILES	CHONES	<none></none>	
		Arabidopsis thaliana					
		cellulose synthase	ļ			1	
		catalytic subunit (Ath-					
		A) mRNA, complete	ł		1		
921	AF027173	cds ·	0.002	<none></none>	<none></none>	<none></none>	
		medium-chain acyl-					
		CoA dehydrogenase			1	1	
		{exon 10, intron 10}		•		l	
000	64107	[human, Genomic,	0.000				
922	S61977	1407 nt]	0.002	<none></none>	<none></none>	<none></none>	
		Arabidopsis thaliana mRNA for					
		neoxanthin cleavage					
923	AJ005813	enzyme	0.002	<none></none>	<none></none>	<none></none>	
		Brassica rapa mRNA	0.002	VI ONE	THE COURT OF THE C	CHOINES	
		for SLG45, complete				l i	
924	AB012105	cds	0.002	<none></none>	<none></none>	<none></none>	
		Brassica rapa mRNA					
		for SRK45, complete					
925	AB012106	cds	0.002	<none></none>	<none></none>	<none></none>	
		A					
ľ		Arabidopsis thaliana cellulose synthase			·	1	
	1	catalytic subunit (Ath-					
		A) mRNA, complete			·	i	
926		cds	0.002	<none></none>	<none></none>	<none></none>	
		H.sapiens DNA for	0.002	ZIONES	(AE001337) Yop C/Gen	MONES	
		dopamine D2	1		Secretion Protein D [Chlamydia		
927		receptor gene	0.002	3329125	trachomatis]	9.5	
				·	HYPOTHETICAL ITZ.T KD		
	•	1	ļ		PROTEIN C06G4.1 IN	!	
ŀ		\$	ļ		CHROMOSOME III	i	
i		1	ŀ		>gi 630524 pir S44748		
j	ļ		ŀ		C06G4.1 protein -		
- 1	1		1		Caenorhabditis elegans		
<u></u>	ť l	İ	j		>gi 409292 (L25598) homology		
),	Mus musculus]		with vigilin; coded for by C. elegans cDNA	İ	
	, .	Pontin52 mRNA.	I		GenBank:M88954 (CEL12C9);		
928		complete cds	0.002	465762	putative [Caenorhabditis	8.9	
		Human skeletal	0.002	403702	pararive [Caenornabultis	0.7	
J		muscle ryanodine			co-repressor protein - mouse	ŀ	
929		receptor gene	0.002	2137221	>gi 642619	6.9	
					1-2-1-2-1		

	Nearest	Neighbor (BlastN vs. (Genhank)	Nassas M	har (DI av	
SEC		Treignoor (Blastia VS.)	Jenoank)	rearest Neigh	nbor (BlastX vs. Non-Redundant P	roteins)
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Mus musculus				
		Pontin52 mRNA,			(Z22520) membrane protein	
930	AF100694	complete cds	0.002	806536	[Bacillus acidopullulyticus]	6.3
		Mus musculus				
		Pontin52 mRNA,	1		(AL023844) Y48A6B.1	
931	AF100694	complete cds	0.002	3881055	[Caenorhabditis elegans]	5.8
932	A F000115	Lycopersicon esculentum cytosolic class II small heat shock protein HCT2 (HSP17.4) mRNA,			(Z81097) K07A1.4	
932	AF090115	complete cds	0.002	3878330	[Caenorhabditis elegans]	4.8
933	AF093268	Rattus norvegicus homer-1c mRNA, complete cds	0.002	137640	REPLICATION PROTEIN E1	4.0
		Mus musculus	0.000	257040	papinomavirus	4.0
		nuclear orphan	ł		(U58757) similar to nucleotide	
934	AF019660	receptor RORgamma	0.002	1330365	pyrophosphatases	3.9
		Mus musculus			(U46951) ORF5; Method:	
		Pontin52 mRNA,			conceptual translation supplied	
935	AF100694	complete cds	0.002	1785972	by author	3.7
		Human gene for			(X56082) protease	
936	V00508	epsilon-globin.	0.002	1333804	[Ruminococcus flavefaciens]	3.5
937	AB012105	Brassica rapa mRNA for SLG45, complete cds	0.002	4153876	(AC005531) similar to mouse homeodomain-interacting protein kinase 2; similar to AF077659 (PID:g3702958)	3.0
938		Arabidopsis thaliana mRNA for neoxanthin cleavage enzyme	0.002	1070461	omithine carbamoyltransferase (EC 2.1.3.3) - yeast (Saccharomyces cerevisiae) >gi 929866 (X83502) pid:c130025 [Saccharomyces cerevisiae] >gi 1008256	2.8
939		rod cGMP phosphodiesterase beta-subunit [human, mRNA, 3231 nt]	0.002	3450883	(AF083334) fibroin [Antheraea	1.6

	<u> </u>	Neare	Nearest Neighbor (BlastN vs. Genbank)		Nearest Ne	ighbor (BlastX vs. Non-Redundant	Dentaire	
	SEQ				1 33.7.0	TEIDER (Blasta Vs. Non-Redundant)	Proteins)	
	ID	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION	N DESCRIPTION		_
			Drosophila			DESCRIPTION .	P VALU	<u></u>
		i	melanogaster Gart					_
	•	Í	locus with genes for				1	
- 1		ſ	GARS=phosphoribo]	i	1	
ſ		ì	ylamineglycine	~	1.			
- 1		ł	ligase,	1	}	i <mark>.</mark>		
- 1			AIRS=phosphoribos	J	j.			
- 1			lformylglycinamidine		ì		1	
- [cyclo-ligase,	-			ĺ	
١			GART=glycinamide		i	1	1	
1			ribotide	1				H
- 1	i		transformylase > ::	-	j		1	1
1			gb J02527 DROGAR	:	[1	1	I
	- 1	'	T D.melanogaster	1	i			ı
-	ſ		Gart gene encoding	1				1
ı	ı	•	two polypeptides with	,] i				1
1			GAR synthase, AIR	Ί			1	1
	- 1		synthase, and GAR]				1
1			transformylase	j			ĺ	ı
	- 1		enzyme activities and]	1
	- 1		a pupal cuticle gene				ļ	1
	- 1		nested within intron				1	I
Ŀ	940	X06286	A of the Gart gene.	0.002	2662054	(A Doored)		ı
Γ			Homo sapiens RNA	0.002	2002034	(AB004651) isocitrate lyase	1.5	1
ı	1		helicase p68	1		1		L
	J		(HUMP68) gene,	l i		(AB008374) alpha 3 type 1		ı
L	941	AF015812	complete cds	0.002	3641659	collagen	l	ŀ
					30 (103)	ZINC FINGER PROTEIN ZFP-	1.1	1
			H.sapiens HZF2			37 (MALE GERM CELL		ı
١.			mRNA for zinc finger	ļ		SPECIFIC ZINC FINGER		L
عل	42	X78925	protein	0.002	141624	PROTEIN)	1.0	
		į					1.0	ı
			Sambucus nigra	1		(Z49071) weak similarity with		1
٥	43	AF074386	hevein-like protein			mu-type opioid receptor (Swiss		ı
	~ }-	AF0/4386	mRNA, complete cds	0.002	3879997	Prot accession number (P33535)	1.0	ı
			Human DNA	j				l
			sequence from			1 .		
			cosmid L241B9,	ł				İ
			Huntington's Disease	ł		. 1		
			Region, chromosome	I		1		
	1		4p16.3 contains			1	j	
			polymorphic VNTR	- 1		LA FORCEON TOTAL		
94	4		pYNZ32.	0.002	3523162	(AF076292) TGF-beta/activin	[
			•		20102	signal transducer FAST-1p	0.81	

Nearest Neighbor (BlastN vs. Genbank) Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		-	Near	est Neighbor (BlastN vs.	Genhank)	Nearest Neighbor (Blast Viva No. 2)			
Date Accession Description Pvalue Accession Description Pvalue Accession Description Pvalue Accession Description Activation Activation Activation Accessi		SE			Genoank)	Nearest Neig	hbor (BlastX vs. Non-Redundant Po	roteins)	
Sambucus nigra hevein-like protein hypothetical protein B - chestnut blight fungus hypothetical protein B - chestnut blight fungus 0.72		4	-	ON DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
Neverin-like protein Neverin-like protein		-							
Martia norvegicus Martia norvegicus Martia norvegicus Martia norvegicus Martia norvegicus Momer- Ic mRNA, complete cds Martia norvegicus Momer- Ic mRNA, complete cds Martia norvegicus Ma		Į							
Section Sect		94	5 A E07430				(AE000761) hypothetical		
AF093268 homer-1c mRNA, complete cds 0.002 101830 hypothetical protein B - chestnut blight fungus 0.72		-	74.07438		s 0.002	2984161	protein [Aquifex aeolicus]	0.80	
AF093268 Complete cds			J						
Homo sapiens Ets-related transcription factor (ERT) mRNA, complete cds 0.002 200531 (M18071) prion protein [Mus musculus] 0.72		94	6 AF09326		0.002	10.000			
Paragraphic Paragraphic				- Jacks CGS	0.002	101830	blight fungus	0.72	
AF017307 AF017307 Complete cds 0.002 200531 (M18071) prion protein [Mus musculus] 0.72]		1	·]	
Drosophila melanogaster Ovo- melanogaster (K80350) Ovo- melanogaster (Manogaster Ovo- melanogaster (Manogaster Ovo- melanogaster (Manogaster Ovo- melanogaster (Manogaster Ovo- melanogaster (Manogaster Ovo- melanogaster (Manogaster Ovo- melanogaster (Manogaster Ovo- melanogaster (Manogaster Ovo- me		1					(2419071)		
Drosophila melanogaster Ovo- 1028a1 (ovo) mRNA, complete cds. 0.002 2465207 (AF016045) OVO-like 1 binding protein [Homo sapiens] 0.35		947	AF017307			200531			
Melanogaster Ovo- 1028aa (ovo) mRNA, complete cds.		1			-	, 200331	musculusj	0.72	
1028aa (ovo) mRNA, complete cds. 0.002 2465207 (AF016045) OVO-like l		ĺ		Drosophila ·					
Sambucus nigra ribosome inactivating protein precursor mRNA, complete cds 0.002 3834294 (U80846) No definition line found (Caenorhabditis elegans) 0.29			1		1	٠			
Sambucus nigra ribosome inactivating protein plant protein plant protein plant protein plant plant protein plant plant protein plant plant protein plant p		l	1	1028aa (ovo) mRNA,	.]		(AF016045) OVO-like 1	i.	
Sambucus nigra ribosome inactivating protein precursor mRNA. complete cds		948	U11383	complete cds.	0.002	2465207		0.35	
Page		i	I				By county (12emo suprems)	0.33	
Saureus genes encoding Sau961 DNA Brassica rapa mRNA Sau961 restriction endonuclease DNA Saureus genes ONO2 Section potential ONO2 Section potential ONO2 Section potential ONO3 ONO2 Section potential ONO3 O			1				1	1	
949 AF012899 mRNA. complete cds 0.002 3834294 found [Caenorhabditis elegans] 0.29			1		1		1.		
Sampa Samp		949	AE012800						
broadening potassium channel=Shab [Aplysia, bag cell neurons, head ganglia. Peptide. 905 and [Aplysia] >gi/34110[prff]2011375A K channel [Aplysia] californica] 0.15 S.aureus genes encoding Sau961 DNA methyltransferase and Sau961 restriction endonuclease 0.002 2529575 3C [Homo sapiens] 0.11 Brassica rapa mRNA for SLG45, complete cds 0.002 729918 AUTOANTIGEN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 G.gallus RAR-gamma2 mRNA for retinoic acid receptor 0.002 586122 TRICHOHYALIN >gi/42321[pirf]/440691 trichohyalin - sheep >gi/295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. mRNA 3231 ml] 0.003 101267		777	A1012899	mikina, complete cds	0.002	3834294	found [Caenorhabditis elegans]	0.29	
Homo sapiens full length insert cDNA clone ZD52F10 0.002 545067 545067 Saureus genes encoding Sau961 DNA methyltransferase and Sau96I restriction endonuclease 0.002 2529575 3C [Homo sapiens] 0.11					i i				
Homo sapiens full length insert cDNA clone ZD52F10 0.002 545067 sqi [Aplysia] >gi 743110[prf] 2011375A K channel [Aplysia californica] 0.15 S.aureus genes encoding Sau961 DNA methyltransferase and Sau96I restriction endonuclease 0.002 2529575 aC [Homo sapiens] 0.11 Brassica rapa mRNA for SLG45, complete cds 0.002 729918 AB012105 Cds 0.002 TRICHOHYALIN >gi 423321[pir] A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 Tod cGMP phosphodiesterase beta-subunit [human. mRNA 3231 nt] 0.003 TATICAS (X90569) elastic titin [Homo	١		1		l			ı	
950 AF086315 AF0	ı		i					i i	
950 AF086315 length insert cDNA clone ZD52F10 0.002 545067 sqif743110 prf 2011375A K channel [Aplysia californica] 0.15	I		ł	Homo sapiens full			neurons, head ganglia, Peptide,	Í	
950 AF086315 clone ZD52F10 0.002 545067 channel [Aplysia californica] 0.15	ļ		ł	1 .	1			ł	
S.aureus genes encoding Sau961 DNA methyltransferase and Sau961 restriction endonuclease 952 AB012105 G.gallus RAR- gamma2 mRNA for retinoic acid receptor 754 S41458 MRNA 3231 ml 954 S41458 S.aureus genes encoding Sau961 DNA methyltransferase and Sau961 restriction 0.002 2529575 (AF018164) kinesin-like protein 3C [Homo sapiens] 0.11 LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 TRICHOHYALIN >gil423321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073	l	950	AF086315		0.002	545067			
encoding Sau961 DNA methyltransferase and Sau961 restriction endonuclease 951 X53096 Brassica rapa mRNA for SLG45, complete cds 0.002 729918 C. Homo sapiens] LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 TRICHOHYALIN 961 S41458 TOT CGMP phosphodiesterase beta-subunit [human. 954 S41458 MRNA 3231 m] 0.002 (AF018164) kinesin-like protein 0.011 (AF018164) kinesin-like protein 0.011 (AF018164) kinesin-like protein 0.011 (AF018164) kinesin-like protein 0.011 (AF018164) kinesin-like protein 0.011 TA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 (X993321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073	ĺ					313007	Chainer [Aprysia californica]	0.15	
DNA methyltransferase and Sau96I restriction endonuclease 0.002 2529575 3C [Homo sapiens] 0.11 Brassica rapa mRNA for SLG45, complete cds 0.002 729918 AB012105 G.gallus RAR- gamma2 mRNA for retinoic acid receptor 0.002 S86122 (AF018164) kinesin-like protein 3C [Homo sapiens] 0.11 LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 TRICHOHYALIN >gi 423321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3731 nt) 0.003 HOLT407	I				1			- 1	
methyltransferase and Sau96I restriction endonuclease 0.002 2529575 3C [Homo sapiens] 0.11 Brassica rapa mRNA for SLG45, complete cds 0.002 729918 LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 G.gallus RAR-gamma2 mRNA for retinoic acid receptor 0.002 586122 TRICHOHYALIN >gi 423321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. mRNA 3231 mt] 0.003 10.07467	ı						1	1	
Sau96I restriction endonuclease 0.002 2529575 (AF018164) kinesin-like protein 3C [Homo sapiens] 0.11 Brassica rapa mRNA for SLG45, complete cds 0.002 729918 LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 G.gallus RAR-gamma2 mRNA for retinoic acid receptor 0.002 586122 TRICHOHYALIN >gi 423321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3231 ml) 0.003 (X90569) elastic titin [Homo	l.		i .	1	ì			I	
951 X53096 endonuclease 0.002 2529575 3C [Homo sapiens] 0.11 Brassica rapa mRNA for SLG45, complete cds 0.002 729918 LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 G.gallus R.AR-gamma2 mRNA for retinoic acid receptor 0.002 586122 TRICHOHYALIN >gi 423321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3231 ml) 0.003 (X90569) elastic titin [Homo	ľ				·	•		ľ	
Brassica rapa mRNA for SLG45, complete cds 0.002 729918 AUTOANTIGEN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 G.gallus RAR-gamma2 mRNA for retinoic acid receptor 0.002 586122 TRICHOHYALIN Sqi 423321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3231 ml) 0.003 INTAGE (X90569) elastic titin [Homo	ı	051	VENDO				(AF018164) kinesin-like protein	1	
952 AB012105 cds 0.002 729918 LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG) 0.092 G.gallus R.AR-gamma2 mRNA for retinoic acid receptor 0.002 586122 TRICHOHYALIN >gi 423321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3231 mt) 0.003 (X90569) elastic titin [Homo	۲	731	A33096		0.002	2529575	3C [Homo sapiens]	0.11	
952 AB012105 cds 0.002 729918 AUTOANTIGEN HOMOLOG) 0.092 G.gallus RAR-gamma2 mRNA for retinoic acid receptor 0.002 586122 TRICHOHYALIN Sqi 423321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3231 ml									
G.gallus RAR- gamma2 mRNA for retinoic acid receptor 0.002 586122 TRICHOHYALIN >gi 423321 pir A40691 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human.		952	AB012105		0.000	5000	RIBONUCLEOPROTEIN) (LA		
953 X73973 G.gallus R.AR- gamma2 mRNA for retinoic acid receptor 0.002 586122 S86122 S86122 (Z18361) trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3231 pt) 0.003 (X90569) elastic titin [Homo			3-2-312103		0.002	/29918	AUTOANTIGEN HOMOLOG)	0.092	
953 X73973 G.gallus R.AR- gamma2 mRNA for retinoic acid receptor 0.002 586122 S86122 S86122 (Z18361) trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3231 pt) 0.003 (X90569) elastic titin [Homo		- 1			-		TRICHOUNTER		
gamma2 mRNA for retinoic acid receptor 0.002 586122 trichohyalin - sheep >gi 295941 (Z18361) trichohyalin 0.073 rod cGMP phosphodiesterase beta-subunit [human.		ł		G.gallus RAR-				- 1	
red cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3231 nt) 965				gamma2 mRNA for	ı				
rod cGMP phosphodiesterase beta-subunit [human. 954 S41458 mRNA 3231 nt) 0.003 (X90569) elastic titin [Homo	_	953	X73973	retinoic acid receptor	0.002	586122	(718361) trickohyalia	0.070	
phosphodiesterase beta-subunit [human. (X90569) elastic titin [Homo		ĺ					(210301) trichonyalin	0.075	
beta-subunit [human.] 954 S41458 mRNA 3231 nt) 0.003 (X90569) elastic titin [Homo		j			ſ]		1	
954 S41458 mRNA 3231 nt) 0.003 (A90309) erastic titin (Homo		- 1	{		ł			- 1	
224 241438 IMKNA (7(17t) 0.000 1.017407 1	•	354			1		(X90569) elastic titin (Homo		
	_	,,,4	341438	mKNA, 3231 nt]	0.002			0.013	

-	Near	est Neighbor (BlastN v	vs. Genbank)	Nearest Nei	ighbor (BlactY ve No - Political	
SE	Q			1,000,050,140,1	ghbor (BlastX vs. Non-Redundant	Proteins)
II.	ACCESSI		N P VALUE	ACCESSION	DESCRIPTION	P VALU
\vdash		D. melanogaster			(U88169) similar to	I VALU
i		defective chorion-			molybdoterin biosynthesis	
١؞		fc 125 (dec-1) gene	e, [MOEB proteins [Caenorhabdit	. [
95	5 M3588	complete cds.	0.002	1825606	elegans]	
				1025000	eregans	0.008
i i	1	Laccaria bicolor		I		1
1	i	glyoxal malate	1	1		- [
۱	. 1	synthase protein		j	(T188167) P2002 2	
950	6 AF03409		ds 0.002	1825593	(U88167) D2092.2 gene produ	1
	1	Bactrocera dorsalis		.023373	[Caenorhabditis elegans]	le-06
	1	strain Tahiti			1	1
	· I	mitochondrial D-lo	ор	1.] .	-
	1 -	region, complete				1
957	AF033929	sequence	9e-04	<none></none>		
		Brassica rapa mRN.		CIVONES	<none></none>	<none></none>
	1	for SRK45, complet				
958	AB012106	cds	8e-04	-NONTE-	f .	
			30.04	<none></none>	, <none></none>	<none></none>
	I	Homo sapiens DEA	D			
	1	box protein (BAT1)			l	1
959	AF029062	gene, partial cds	8e-04	-NONE-		
		Human ataxin-2	30-04	<none></none>	<none></none>	<none></none>
		related protein				
960	U70671	mRNA, partial cds	8e-04	<none></none>		
		Dendrocopos	1 30-07	CIVOINES	<none></none>	<none></none>
	l	leucopterus clone 2	1 1			
	ļ	microsatellite HrU2	1		•	
961	AF051709	repeat region	8e-04	NONE		
		Pea phy gene for	06-04	<none></none>	<none></none>	<none></none>
		phytochrome	1 1			
962	X14077	apoprotein	8e-04	-NONT		ı İ
		Homo sapiens	00-04	<none></none>	<none></none>	<none></none>
ı		chromosome 21, P1	1 1			
63	AC004497	clone LBNL#6	8e-04	467144	(L27838) rhoptry protein	. [
T		Homo sapiens	30-04	457146	[Plasmodium yoelii]	9.6
J		cartilage-derived C-	 			
64	AF077344	type lectin	8004	2702/22	(AJ011707) TraD protein	ĺ
		77- 100	8e-04	3702123	[Escherichia coli]	8.5
		H.sapiens epb72 gene	1 1		(AJ004687) N-4 cytosine-	
65		exons 2.3.4,5,6,7			specificmMethyltransferase	- 1.
			8e-04	2570059	[Neisseria gonorrhoeae]	6.8
- 1		ļ	1	I		
- 1]		1	<u> </u>	COPPER TRANSPORT	ı
	I	Mus musculus	ĺ	 1	PROTEIN CTR transport	· · ·]
		Pontin52 mRNA,	1	ļī	protein - yeast (Saccharomyces	1
6		complete cds	9- 0-	la	cerevisiae) gene product	j
		-ombiere cas	8e-04		Saccharomyces cerevisiae]	

<u> </u>	: Near	est Neighbor (BlastN vs.	Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant)	
	EQ				(Blasta Vs. Non-Redundant	Proteins)
	D ACCESSI		P VALUE	ACCESSION	DESCRIPTION	
_		Homo sapiens			DESCRIPTION	P VALU
	1	MLL/AF4			SMALL PROTEIN B	
	j	translocation		1	HOMOLOG A43259, from E.	
96	7 AF03140	breakpoint 13 t(4;11)(q21;23)	1		hirae [Mycoplasma	
	12 03140	Human (clone D13-2	8e-04	2498926	pneumoniae)	6.6
1	ſ	L-iditol-2	~1			
	1	dehydrogenase gene,		1	İ	
1	1	exon 4, exon 5, exon		1	(1163007)	
96	8 L29252	6 and exon 7.	8c-04	1488070	(U63997) putative transposase [Enterococcus faecium]	
	1	Mouse N10 gene for			(Enterococcus faecium)	5.2
969	V16005	a nuclear hormonal	1	•	(U47323) stromal cell protein	
1	X16995	binding receptor	8c-04	1493833	[Mus musculus]	3.2
	1	i	1			
1	ł				4-AMINOBUTYRATE	ł
ļ	1	j			AMINOTRANSFERASE	
l	1		1 1		TRANSAMINASE) (GABA	1
	1				AMINOTRANSFERASE) homolog - smut fungus]
	j		1		(Ustilago maydis) >gi 881562	1
l	Ĭ	<u> </u>	1 1		Emericella nidulans gamma-	[
	1	Human interleukin-8	1		amino-n-butyrate transaminase	
970	M99412	receptor (IL8RB)			Swiss-Prot Accession Number	
770	14199412	gene, complete cds Human Down	8e-04	1346101	P14010 [Ustilago maydis]	0.83
		Syndrome region of	1			
	1	chromosome 21	1		1	i
		genomic sequence,			(1E114000)	1
971	U37452	clone A31D6-1C5.	8e-04	4164069	(AF111093) latrophilin 3 splice	1
	1			4104009	variant bbah [Bos taurus] HYPOTHETICAL 13.0 KD	0.26
	1		ſ		PROTEIN IN RAD26-GEF1	
		1		ĺ	INTERGENIC REGION	ı
	•				>gi 1077881 pir \$57057	
		1.]	probable membrane protein	
			j	l l	YJR038c - yeast	
		Mus musculus	į	19	(Saccharomyces cerevisiae)	
		Pontin52 mRNA.	. 1	Į:	>gi 1015688 (Z49538) ORF	- 1
972	AF100694	complete cds	8e-04	1252022	YJR038c putative	
T		Rattus norvegicus	30-04	1352877	Saccharomyces cerevisiae]	0.23
[homer-1c mRNA,	I	1,	A F000312) ==6 11	
973	AF093268	complete cds	8e-04		AE000312) orf, hypothetical protein [Escherichia coli]	ł

	Nearest	Neighbor (BlastN vs. C	ienbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC		1	T	realest Neigh	ibor (Blasta Vs. Non-Redundant P	roteins)	
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
-		 			HYPOTHETICAL 37.7 KD		
1	1		1		PROTEIN C18B11.06 IN		
1	ł				CHROMOSOME I	j	
	1		Ĭ,		>gi 2130289 pir S58305	1	
1	1	H.vulgaris mRNA for	1		hypothetical protein	I	
ł		cAMP response	1		SPAC18B11.06 - fission yeast		
	ľ	element binding	<u> </u>		hypothetical protein	ł	
974	X83872	protein	8e-04	1175386	[Schizosaccharomyces pombe]	0.005	
						-	
075	1,50051.	Rat simple sequence			(AF024502) No definition line	1	
975	M32514	DNA, clone 5.	8e-04	2394492	found [Caenorhabditis elegans]	0.002	
ł		Sambucus nigra	٠.				
ļ		hevein-like protein			(AB012223) ORF2 [Canis	1	
976	AF074386	mRNA, complete cds	8e-04	2981631	familiaris]	0.001	
						0.001	
]	1	H.sapiens DNA for	İ]	
077	******	endogenous retroviral			(Y12713) Pro-Pol-dUTPase	1	
977	X89211	like clement	8e-04	2065210	polyprotein	3e-04	
ļ		,			(AC002411) Strong similarity to		
		Human myosin-IC	i		myosin heavy chain gb Z34293 from A. thaliana. [Arabidopsis		
978	U14391	mRNA, complete cds.	. 8e-04	3142302	thaliana)	4e-16	
		Drosophila		3112302	tiaiai	46-10	
]		melanogaster dead-				l l	
		box protein					
		D.melanogaster	1		*		
979	1.12610	DEAD-box gene,			(AJ010475) RNA helicase		
7/7	L13612	complete CDS	. 8e-04	3776027	[Arabidopsis thaliana]	9e-24	
		Sambucus nigra	[1		
		hevein-like protein	i				
980	AF074386	mRNA, complete cds	7e-04	<none></none>	<none></none>	<none></none>	
		Mus musculus					
		Pontin52 mRNA,	f		1	ł	
981		complete cds	7e-04	<none></none>	<none></none>	<none></none>	
		Rattus norvegicus homer-1c mRNA,	1				
982		complete cds	7e-04	<none></none>	NONE		
		Human DNA	70-04	CHOINES	<none></none>	<none></none>	
		sequence from	1			j	
1	ŀ	cosmid N120B6 on	1	ļ		i	
į		chromosome 22	f			i	
		Contains ESTs,	l l	ł		Į	
202		complete sequence					
983	Z73987	Homo sapiens]	7e-04	<none></none>	<none></none>	<none></none>	

	Neares	t Neighbor (BlastN vs. (Genbank)	Nearest Nai	ghbor (BlastX vs. Non-Redundant	
SE			T	TVCAICST IVE	gilbor (BlastX Vs. Non-Redundant	Proteins)
ID	-	N DESCRIPTION	PVALUE	ACCESSION		
		Brassica rapa mRNA		ACCESSION	DESCRIPTION	P VALUE
		for SRK45, complete				
984	AB012106		7e-04	<none></none>		
		Rattus norvegicus	1 70 07	KINONES	<none></none>	<none></none>
	į.	homer-1c mRNA,	1			ĺ
985	AF093268	complete cds	7e-04	<none></none>	<none></none>	
				4.0	CHONES	<none></none>
	1	Arabidopsis thaliana			1	
ł	1	cellulose synthase	Į l		· [1
1		catalytic subunit (Ath-	1		.	
986	45005151	B) mRNA, complete			1	
980	AF027174		7e-04	<none></none>	<none></none>	<none></none>
ł		Mus musculus	٠. ا			
987	AF100694	Pontin52 mRNA, complete cds	7 04			
1	AL 100094	Arabidopsis thaliana	7e-04	<none></none>	<none></none>	<none></none>
	1	mRNA for				
		neoxanthin cleavage				1 1
988	AJ005813	enzyme	7e-04	<none></none>		1 1
			70 07	CHOINES	<none></none>	<none></none>
1		Helianthus tuberosus	1			1
		lectin 1 mRNA,	i			1 1
989	AF064029	complete cds	7e-04	<none></none>	<none></none>	<none></none>
						14.02
	ĺ	Arabidopsis thaliana	1]
	ł .	cellulose synthase			•	1
		catalytic subunit (Ath-B) mRNA, complete				1 1
990	AF027174	cds	7. 04			
		cus	7e-04	<none></none>	<none></none>	<none></none>
		Arabidopsis thaliana	1		1	1
		cellulose synthase	1			i i
		catalytic subunit (Ath-		•	}	1 1
	· /	A) mRNA, complete	1			1
991	AF027173	cds	7e-04	<none></none>	<none></none>	<none></none>
						THE PARTY OF THE P
		Helianthus tuberosus	I]]
992		lectin 1 mRNA,]]
774	AF064029	complete cds	7e-04	<none></none>	. <none></none>	<none></none>
		Mus musculus	1			
993		Pontin52 mRNA,	70.04			
	-11 100074	complete cds	7e-04	<none></none>	<none></none>	<none></none>
İ	l _s	Sambucus nigra	1			
ł		ribosome inactivating	Į			1
- 1		protein precursor	1		(A BO14608) KIA 40703	1
994		nRNA, complete cds	7e-04	3327230	(AB014608) KIAA0708 protein [Homo sapiens]	0.5
				2241230	Itanio sapiens)	9.5

	Nearest	Neighbor (BlastN vs. C	Jenhank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		Reighbor (Blash Vs. C	Jenoank)	Nearest Neigh	nbor (BlastX vs. Non-Redundant Pr	oteins)	
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
<u> </u>							
İ	1	Sambucus nigra					
	1	ribosome inactivating	1				
995	U76524	protein precursor mRNA, complete cds	7 04		(AB014608) KIAA0708 protein		
1 333	070324	mkivA, complete cas	7e-04	3327230	[Homo sapiens]	9.3	
1	l	Sambucus nigra]		(Z93380) predicted using Genefinder; similar to 7tm		
1		hevein-like protein			receptor protein [Caenorhabditis		
996	AF074387	mRNA. complete cds	7e-04	3876455	lelegans?		
			75 57	3070433	hypothetical protein MJ1293 -	7.1	
1				0	Methanococcus jannaschii		
1		Sambucus nigra		٠,	>gi 1591931 (U67570) M.		
		ribosome inactivating			jannaschii predicted coding		
1.		protein precursor			region MJ1293 [Methanococcus		
997	U76524	mRNA, complete cds	7e-04	2128771	jannaschii)	6.2	
			•				
i		Human zinc finger					
998	1100412	protein ZNF134			glutathione transferase (EC	i	
390	U09412	mRNA, complete cds	7e-04	1083336	2.5.1.18) piA - mouse	5.4	
i i		Arabidopsis thaliana					
		cellulose synthase			1		
		catalytic subunit (Ath-			(M17619) NADH	l l	
		A) mRNA, complete	1		dehydrogenase subunit ND4		
999	AF027173	cds	7 c -04	473515	[Asterina pectinifera]	3.7	
i T					[isterna peetimeta)		
i j		Sambucus nigra	j		1		
		ribosome inactivating			1	l	
1000	4 F0 10 00 0	protein precursor	i		(U79772) female sex protein	1	
1000	AF012899	mRNA, complete cds	7e-04	1724097	[Mercurialis annua]	3.3	
- 1		Mus musculus					
1001		Pontin52 mRNA, complete cds	70.04	110710-	(D49747) core, env, and part of	_ }	
1001	AI 100034	complete cas	7e-04	1197103	E2/NS1 unc-5 protein, long form -	3.2	
ĺ			ì		Caenorhabditis elegans	İ	
ł	i		ł		>gi 258529 bbs 118648		
ł	ı	i	ŀ		(S47168) UNC-		
- 1	l	į	ļ		5=immunoglobulin and		
	j	j			thrombospondin type 1	i	
	[ļ			transmembrane protein		
1	j				{alternatively spliced} aa]		
- 1		Mouse N10 gene for			[Caenorhabditis elegans]	1	
		nuclear hormonal	į.		>gi 2662596 (AF036698) C.	1	
1002	X16995 I	oinding receptor	7 c -04	345372	elegans UNC-5 (NID:g25852)	2.7	

	Nearest	Neighbor (BlastN vs. C	ienbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ					I Reduidant	Totems	
ID	ACCESSIO	DESCRIPTION .	P VALUE	ACCESSION	DESCRIPTION	P VALUI	
	<u> </u>						
	j	Sambucus nigra					
1	i	ribosome inactivating	ł	ł	ĺ	1	
1003	U76524	protein precursor			(AB022866) mobilization		
1003	070324	mRNA, complete cds Rattus norvegicus	7e-04	4204220	protein	2.5	
		homer-ic mRNA,					
1004	AF093268	complete cds	70.04	2201550	(Y17116) fibrinogen-binding	1	
1007	712 055200	complete cus	7e-04	3201550	protein	2.4	
	ł	Sambucus nigra					
		hevein-like protein			(1145066) 1	i	
1005	AF074386	mRNA, complete cds	7e-04	1174264	(U45966) polyprotein [Hepatitis		
		and any complete cas	76-04	11/4204	(G virus)	0.73	
		Arabidopsis thaliana	•	l	1		
		cellulose synthase				1	
		catalytic subunit (Ath-	I		i	l	
		A) mRNA, complete	ľ		TRANSCRIPTION FACTOR	ł	
1006	AF027173	cds	7e-04	135308	JUN-D	0.065	
1		H.sapiens EWS gene,				0.003	
		intron 6,	1		!!!! ALU SUBFAMILY SP	1	
1007	X98745	polymorphism	7e-04	728836	WARNING ENTRY	0.001	
1		Arabidopsis thaliana					
- 1		mRNA for	1			1	
1000	4 T005010	neoxanthin cleavage					
1008	AJ005813	enzyme	7e-04	1633564	(U47924) C8 [Homo sapiens]	9e-09	
		Sambucus nigra	1		•		
J		hevein-like protein	1				
1009	AF074386	mRNA, complete cds	6e-04	284171	Ig epsilon chain C region form 3		
		Brassica rapa mRNA	06-04	2041/1	- human (AE001414) BRAHMA	1.3	
- 1		for SRK45, complete			ortholog (DNA helicase		
1010		cds	6e-04	3845262	superfamily II)	0.35	
		Human DNA		30,3202	supertaining II)	0.25	
ı	1	sequence from clone	ł	•			
		417C12 on	- 1			j	
	ŀ	chromosome Xp22.11	i			1	
		22.2, complete	i			i	
	ļ:	sequence [Homo	ł				
011		sapiens]	3e-04	<none></none>	<none></none>	<none></none>	
- 1							
		Iomo sapiens (pp21)	i				
012	M99701 r	nRNA, complete cds.	3e-04	<none></none>	<none></none>	<none></none>	

<u> </u>		st Neighbor (BlastN vs.	Genbank)	Nearest Nei	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SE	- 1	1.				Totems		
II	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU		
\vdash		Ovis aries Merino						
		breed DR beta-chain	1					
i		antigen binding	1	1		l		
1		domain, MHC class	,,					
	ľ	DRB (Ovar-DRB24)				1		
101	3. U00227	gene, partial cds.	3e-04) //O	•			
		gene, partial cus.	Je-04	<none></none>	<none></none>	<none></none>		
	1	Sambucus nigra	1		·			
- 1	1	hevein-like protein	1			1		
1014	AF074387	mRNA, complete cds	3e-04	<none></none>	NONT	1		
				··	<none></none>	<none></none>		
İ	1	Xenopus laevis	j i		f			
ĺ	ł	mitotic			1	1		
		phosphoprotein 90			(L19655) ORF [Tomato	ł		
1015	U95102	mRNA, complete cds	3e-04	999418	ringspot virus]	8.3		
		Brassica rapa mRNA			(AF011415) putative	0.3		
1016	4.0010106	for SRK45, complete	i I		pheromone receptor [Mus	· ·		
1010	AB012106	cds	3e-04	2367460	musculus]	7.0		
		Mus musculus DNA						
		for microsatellite 3kb	1			l i		
1017	AJ010737		2 04		(AF104411) neuronal-specific			
	1.5010757	upstream lbp gene	3e-04	4106549	septin 3 [Mus musculus]	5.5		
	Ī				NADH-DEPENDENT FLAVIN			
1		Homo sapiens histone	ı		OXIDOREDUCTASE acid-			
1		deacetylase 3 gene.			inducible - Eubacterium sp			
1		exons 4, 5, 6, 7, 8, 9,	1	-	>gi 1381570 (U57489)			
1018	AF053137	and 10	3e-04	416702	NADH:flavin oxidoreductase	1		
				410702	[Eubacterium sp. VPI 12708]	5.3		
1		Arabidopsis thaliana			1			
i i		cellulose synthase	ı		!	ł		
		catalytic subunit (Ath-	1	. •		1		
ا		A) mRNA, complete	. [(Y08502) orf111d [Arabidopsis	- 1		
1019		cds	3e-04	1785789	thaliana]	5.1		
		Homo sapiens clone						
	i.	UWGC:y23x011	1]	1		
		from 6p21, complete	- 1		1			
1020		sequence [Homo			(D28917) polyprotein [Hepatitis	ŀ		
1020		sapiens] Human IGF-I mRNA	3e-04	558521	C virus]	1.1		
ĺ		for insulin-like	1					
1021		growth factor I	3- 04	10065:-	(AF118122) putative outer	i		
		FOARI INCIDI I	3e-04	4206707	membrane protein OmpU	0.65		
ı	l _F	Lsapiens IL-IRa	ŀ		(1110700) = 10.05 =			
1022	**	tene.	3e-04	1065941	(U40799) F42C5.7 gene product			
				1003941	[Caenorhabditis elegans]	0.12		

11/2 1	Nearest Neighbor (BlastN vs. Genbank)		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ				ivearest iveign	iooi (DiastA vs. Non-Redundant P	roteins)
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
<u> </u>		Pseudorabies virus				
1		with upstream and			(AF040650) contains similarity	
1	1	downsteam	1		to sodium-potassium-chloride	
1023	M34651	sequences.	3e-04	2746853	cotransport proteins	7e-05
	l				PROBABLE ATP-	1
1		İ]	l .	DEPENDENT RNA	
			1	Ì	HELICASE HAS I	1
1		S.cerevisiae]		>gi 626265 pir S47451	1
i		chromosome II			hypothetical protein YMR290c	1 1
1024	726011	reading frame ORF			RNA helicase [Saccharomyces	
1024	Z36011	YBR142w Dictyostelium	3e-04	2500537	cerevisiae]	4e-08
1		discoideum 2034				
1025	AF020286		ا نو م		(U64857) No definition line	
1025	AI 020286	gene, partial cds	3e-04	1465834	found [Caenorhabditis elegans]	6e-14
İ		Chlamydomonas				
1		reinhardtii dynein				1 1
]		heavy chain alpha			(781077) and inted	
1		(ODA11) gene, exons			(Z81077) predicted using Genefinder; Similarity to Yeast	
1026	L26049	2-15, and partial cds.	3e-04	3876775	protein 8248 (TR:G587531)	0.15
		Dictyostelium		3010113	protein 6248 (TR.C587531)	9e-15
ł I		discoideum 2034			(U64857) No definition line	i .
1027	AF020286	gene, partial cds	3e-04	1465834	found [Caenorhabditis elegans]	le-17
					(Z69635) Similarity to Yeast	10 17
			ľ		uridine kinase	
1 1		ļ			(SW:URKI_YEAST); cDNA	
					EST EMBL:Z14695 comes	1
					from this gene; cDNA EST	
ł		į	1		CEMSE17F comes from this	
			i		gene; cDNA EST	
			ľ		EMBL:D67355 comes from this	
1028	J	S.cerevisiae ACT3			gene; cDNA EST yk209h1.5	1
1028	X79811	gene	3e-04	3876090	comes from this ge	7e-31
		Arabidopsis thaliana	ľ		·	.[
ł		cellulose synthase	1]	ł
ł		catalytic subunit (Ath-	į			
1		A) mRNA, complete	1			- 1
1029		eds	2e-04	<none></none>	NONE	
		Human pancreatic	20-04	SHORES	<none></none>	<none></none>
j		phospholipase A-2			· I	i
- 1		PLA-2) gene, exons	1	•	1	1
1030		L to 3.	2e-04	<none></none>	<none></none>	<none></none>
					~	71.0117

-		Neighbor (BlastN vs.	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC	. 1						
ID	ACCESSIO		P VALUE	ACCESSION	DESCRIPTION	P VALU	
<u> </u>		Human DNA	1			1. VALU	
1		sequence from					
İ	ľ	cosmid N2E9 on	l	i)	i	
1	i	chromosome 22.]		
1	-	Contains EST,			1		
1	1	complete sequence			1	1	
1031	Z68686	[Homo sapiens]	2e-04	<none></none>	<none></none>	27025	
		H.sapiens brca2 gene		4.01.02	CHONES	<none></none>	
1	Ī	exon 4 > ::					
	1	emb A62779 A62779	i				
		Sequence 20 from	ł			1	
1032	X95154	Patent WO9719110	2e-04	·· <none></none>	<none></none>	A TONT	
		Arabidopsis thaliana			CNOINES	<none></none>	
Ì		mRNA for					
		neoxanthin cleavage				Į	
1033	AJ005813	enzyme	2e-04	<none></none>	<none></none>	-NONTE:	
i		Mus musculus			CIVOINES	<none></none>	
i		Pontin52 mRNA,	1				
1034	AF100694	complete cds	2e-04	<none></none>	<none></none>	NONTE	
		Plasmodium				<none></none>	
		falciparum					
		chromosome 2,				i	
		section 52 of 73 of	j			·	
		the complete					
1035	AE001415	sequence	2e-04	<none></none>	<none></none>	<none></none>	
		Lycopersicon				1 1	
		esculentum cytosolic	J		ł	1 1	
		class II small heat	1			[]	
1		shock protein HCT2	į.		1]	
1036		(HSP17.4) mRNA,			· ~	1 1	
1030	AF090115	complete cds	2e-04	<none></none>	<none></none>	<none></none>	
l		Homo sapiens (subclone 6_d9 from	. [
ł		PI H21) DNA	į	•] [
1037		sequence]	
1037		Rattus norvegicus	2e-04	<none></none>	<none></none>	<none></none>	
		nomer-1c mRNA,	1.				
1038		, ,	2.04		CD59 GLYCOPROTEIN		
- 320	1093200	complete cds	2e-04	2501523	PRECURSOR	7.1	
	ls	Sambucus nigra	- 1				
		ibosome inactivating	1			1	
		rotein precursor	ı			1	
1039		RNA, complete cds	2e-04	2765262	(Y13925) cathepsin L2 [Penacus	į.	
		complete cus [≟C+U4	2765360	vannameil	6.8	

	Nearest	Neighbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC	2				- Control of the Cont	Totellis,	
ΙD	-	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUI	
<u> </u>					RNA POLYMERASE		
1		Arabidopsis thaliana			>gi 67126 pir RRXPLC RNA-		
1		cellulose synthuse	1	1	directed RNA polymerase (EC		
1	1	catalytic subunit (Ath	1	Í	2.7.7.48) - lymphocytic	ĺ	
		B) mRNA, complete		ł	choriomeningitis virus (strain	ł	
1040	AF027174	cds	2e-04	133636	. Armstrong 53b) >gi 331369	5.2	
1	İ	Brassica rapa mRNA			(AF074613) type II secretion		
1		for SRK45, complete			protein [Escherichia coli	1	
1041	AB012106	cds	2e-04	3822155	O157:H7]	4.0	
			1				
1		Sambucus nigra				1	
i .		ribosome inactivating		• •	-	1	
1,040	7,70,000	protein precursor			REGULATORY PROTEIN E2		
1042	U76524	mRNA, complete cds	2e-04	1718125	>gi 1020222 type 36]	0.38	
1	į	Sus scrofa mRNA for					
1043	3/15050	glucose transporte				1	
1043	X17058	protein	2e-04	3341906	(AB009593) xylose transporter	2e-15	
		Homo sapiens					
1044	A.E.O.O.O.	candidate tumor					
1044	AF008216	suppressor pp32r1	le-04	<none></none>	<none></none>	<none></none>	
		S.tuberosum mRNA					
l	l	for inorganic	ľ			1	
1045	X98890	phosphate			(U42580) a65L [Paramecium]	
1043	798890	transporter, StPT1	le-04	624126	bursaria Chlorella virus 1]	7.9	
		Chaire (D. La.)	l				
1046	L14930	Glycine max (Rab7p)					
1040	L14930	mRNA, complete cds. Mus musculus	9e-05	<none></none>	<none></none>	<none></none>	
		thromboxane A2	Į.			1	
						i :	
1047		receptor gene, exon 3,	0.05				
1047		M.musculus mRNA	9e-05	<none></none>	<none></none>	<none></none>	
1048		for Brx gene, partial	0- 05	\	1		
-0.00	111890	ioi bix gene, partiai	9e-05	<none></none>	<none></none>	<none></none>	
į	ŀ	Polistes annularis			·		
ı		(clone pan48AAT)	i		l l		
1049		andem repeat region.	9e-05	ANONIT-			
		unden repeat region.	90.03	<none></none>	<none> (Z79758) cDNA EST</none>	<none></none>	
- 1	j		1		EMBL:D28009 comes from this		
ŀ					gene; cDNA EST	Ţ.	
			1		EMBL:D28008 comes from this	1	
]	1		i		gene; cDNA EST		
- 1	1		1		EMBL:D32478 comes from this	i	
- [l.		j		gene; cDNA EST	ł	
- 1	1	1				i	
- 1	1.	Iomo sapiens clone	j		EMBL:D34508 comes from this	f	
		4587 mRNA	.1		gene; cDNA EST	1	
1050		equence	9e-05	3880586	EMBL:D37581 comes from this	-,	
			70.03	000000	gene;	7.6	

	Nearest	Neighbor (BlastN vs. C	ienbank)	Negreet Neig	hhos (Bloov)	
SEQ		T	T T	i vearest iverg	hbor (BlastX vs. Non-Redundant P	roteins)
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-	 	Cambana				
1051	U76524	Sambucus nigra ribosome inactivating protein precursor mRNA, complete cds	9e-05	3024292	RHODOPSIN >gi 2290717 (AF000947) rhodopsin [Sepia officinalis]	6.7
1052	Z58294	H.sapiens CpG DNA, clone 34d6, forward read cpg34d6.ftla	9e-05	3885496	(AF064825) heparin/heparan sulfate N-acetylglucosaminyl N-deacetylase/N-sulfotransferase [Bos taurus]	
1053	D87451	Human mRNA for KIAA0262 gene, complete cds	9e-05	3874739	(Z66495) similar to claustrin like	0.004
1054	L37092	Mus musculus cyclin- dependent kinase homologue	 9e-05	3080513	(AL022598) hypothetical protein	4e-09
1055	AF074386	Sambucus nigra hevein-like protein mRNA, complete cds	8e-05	<none></none>	<none></none>	<none></none>
1056		Arabidopsis thaliana cellulose synthase catalytic subunit (Ath-B) mRNA, complete	8e-05	NONE		
1057		Sambucus nigra hevein-like protein mRNA, complete cds	8e-05	<none></none>	<none></none>	<none></none>
1058		Homo sapiens DNA from cosmid clone:844, GT repeat	8e-05	<none></none>	<none></none>	<none></none>
1059	e s p	-ycopersicon sculentum class II mall heat shock protein Le-HSP17.6	0.05		HYPOTHETICAL 80.4 KD PROTEIN IN SMC3-MRPL8 INTERGENIC REGION >gi 1078237 pir \$56849 probable membrane protein YJL073w - yeast (Saccharomyces cerevisiae) >gi 895898 (X88851) hypothetical protein YJL073w	
	ŀ	I. sapiens XB gene	8e-05		[Saccharomyces cerevisiae] microtubule-associated protein, 110K tau - rat >gi 207158	6.0
060		or tenascin-X, repeat	8e-05		(M84156) big tau [Rattus: norvegicus]	3.7

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (Blast X vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
1061	AF027174	Arabidopsis thaliana cellulose synthase catalytic subunit (Ath-B) mRNA, complete cds	8e-05	4049682	(AF063866) ORF MSV092 hypothetical protein [Melanoplus sanguinipes entomopoxvirus]	2.1	
		Lycopersicon esculentum cytosolic class II small heat shock protein HCT2 (HSP17.4) mRNA,		-	(AJ235271) unknown		
1062	AF090115	complete cds	8e-05	3861019	[Rickettsia prowazekii]	5e-14	
		Arabidopsis thaliana cellulose synthase catalytic subunit (Ath-B) mRNA, complete					
1063	AF027174	cds	7e-05	<none></none>	<none></none>	<none></none>	
1064	L04193	Human lens membrane protein (mp19) gene, exon	2. 05	No.			
1004	L04193	11.	7e-05	<none></none>	<none></none>	<none></none>	
1065		B.napus gene for LHC II Type III chlorophyll a/b binding protein	7e-05	2132314	yeast similarity to a nuclear lamin from C. elegans (PIR accession number S42257) [Saccharomyces cerevisiae]	8.9	
		January Protein		2132314	[Oaccharonives cerevisiae]	6.5	
1066		Helianthus tuberosus lectin 1 mRNA, complete cds	7e-05	2979422	(AB006757) PCDH7 (BH- Pcdh)c [Homo sapiens]	5.7	
	6	Arabidopsis thaliana cellulose synthase catalytic subunit (Ath-A) mRNA, complete	·		HYPOTHETICAL 21.5 KD PROTEIN (ORF 185) >gi 1480440 (U34204) ORF185; hypothetical 21.4 kD		
1067	AF027173 d	cds	7e-05	2493696	protein [Brassica oleracea]	5.2	
	B	Rattus norvegicus			PROBABLE LEUCYL-TRNA SYNTHETASE, MITOCHONDRIAL PRECURSOR (LEUCINE TRNA LIGASE) (LEURS)		
1068	AF093268	complete cds	7e-05	2501029	KIAA0028 [Homo sapiens]	1.4	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ				(Mealest Neighbor (Blastx Vs. Non-Redundant Proteins)			
ID	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Human DNA					
i		sequence from				 	
1		cosmid cN85E10 on		j		ĺ	
		chromosome 22q11.2-	1				
1069	Z68758	qter	3e-05	<none></none>	<none></none>	<none></none>	
		human Histone H3.3					
		pseudogene (CIR-					
1070	X60653	456)	3e-05	<none></none>	<none></none>	<none></none>	
]	·	Warri C C DNA					
1		H.sapiens CpG DNA,			GUANYLYL CYCLASE GC-E		
1071	Z58294	clone 34d6, forward	0.05		PRECURSOR cyclase receptor		
10/1	238294	read cpg34d6.ft1a. Homo sapiens	3e-05	1706241	[Mus musculus]	9.6	
		mitochondrial outer			1		
		membrane protein		,	}		
		(Tom40) gene,			AMINE OVIDAGE IEL AVDI		
1 1		nuclear gene			AMINE OXIDASE (FLAVIN-		
1 1		encoding			CONTAINING] B oxidase		
1 1		mitochondrial			(flavin-containing) (EC 1.4.3.4) B - human B [human, platelet,		
		protein, exons 1					
1072	AF043251	through 6	3e-05	113980	Peptide Partial, 520 aa] [Homo sapiens]		
			30.03	113980	sapiens	8.9	
[1	- 1	•		ł	
i 1		Chicken progesterone				i	
! !		receptor gene,				i	
		encoding forms A and			IG GAMMA LAMBDA	ŀ	
1073	M31104	B. exons 1 and 2.	3e-05	1170841	CHAIN V-II REGION	4.8	
			1			•	
		Sambucus nigra	ĺ			1	
j	ł	ribosome inactivating	l l		ribosomal protein S3 -	I	
1074	15010000	protein precursor			Chlamydomonas humicola		
1074		mRNA, complete cds	3e-05	543684	chloroplast (fragment)	4.2	
	1	Human vasopressin receptor V2 gene,	ł		4120616		
i075		complete cds.	30.05	701207	(U20615) Gnot1 homeodomain		
-0/5		Rattus norvegicus	3e-05	791207	protein [Gallus gallus]	1.8	
		homer-1c mRNA,	l		(AE032361) polymentin	1	
1076	1	complete cds	3e-05		(AF033361) polyprotein [Hepatitis C virus]	0.04	
1		Mus musculus	30-03	3631340	[mepatitis C virus]	0.94	
- 1	li li	Pontin52 mRNA,	·		(AL021813) hypothetical	ì	
1077		complete cds	3e-05		protein	0.001	
		Mus musculus				0.001	
ŀ	1	Pontin52 mRNA,	1	i,	(Z81555) predicted using	}	
1078		complete cds	3e-05		Genefinder	3e-07	

<u> </u>	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC						T	
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
-		Lycopersicon					
		esculentum cytosolic					
1	ł	class II small heat	!				
1		shock protein HCT2			·	1	
		(HSP17.4) mRNA.				· ·	
1079	AF090115	complete cds .	2e-05	<none></none>	<none></none>	<none></none>	
	l	77 11 11 11					
	J	Helianthus tuberosus lectin 1 mRNA,				İ	
1080	AF064029	complete cds	2e-05	200107	(Z81132) predicted using	, ,	
1	111 00 1027	Homo sapiens full	26-03	3880197	Genefinder	2.4 .	
		length insert cDNA			!!!! ALU CLASS B WARNING		
1081	AF087989	clone YX29D10	2e-05	113667	ENTRY !!!!	1.8	
1	[Helianthus tuberosus	ı			}	
1082	A F0< 4000	lectin 1 mRNA,			(L31967) mating type protein		
1082	AF064029	complete cds	2e-05	474896	[Coprinus cinereus]	1.4	
	l	Helianthus tuberosus	I				
		lectin 1 mRNA,	ı		(Y13274) M33 polycomb-like		
1083	AF064029	complete cds	2e-05	2266988	protein [Mus musculus]	0.62	
		Equus caballus UCD-			[0.02	
l		E-CA-467					
ļ		dinucleotide repeat	I				
1084	U67415	region, complete					
1004	00/413	sequence H.sapiens BGP gene	1e-05	<none></none>	<none></none>	<none></none>	
		for biliary	ł				
	/	glycoprotein,	Į.				
		promoter region and	J		· ·		
1085	X67277	exon 1	1e-05	<none></none>	<none></none>	<none></none>	
					·		
1086		H.sapiens epb72 gene	1.05				
1080	X85117	exons 2,3.4,5,6,7	1e-05	<none></none>	<none></none>	<none></none>	
	t	Mus musculus	1		(Z29457) core region;		
l		suppressor of			pid:g443877 [Hepatitis C virus]		
1087		cytokine signalling-3	1e-05	443877	virus]	3.9	
					(Z66498) similar to cuticle		
į			- 1		collagen; cDNA EST		
1088		Homo sapiens P2X7			EMBL:D75584 comes from this	İ	
1088	Y12853	gene, exon 4-8	1e-05	3878726	gene	0.36	
- 1		1	ł		(AJ235270) GLUTAMYL-		
ļ	İı	Borrelia burgdorferi	ı		RNA AMIDOTRANSFERASE		
]		section 26 of 70) of	1		SUBUNIT A (gatA) [Rickettsia		
1089	AE001140 t	he complete genome	1e-05	3860719	prowazekii)	4e-15	

	Nearest Neighbor (BlastN vs. Genbank)			Negroes N	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEO		10.00.00.00.00.00	T T	rvediest iveigi	nbor (Blasta vs. Non-Redundant P	roteins)		
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
ļ						T		
1	ł	Homo sapiens gamma				†		
		adaptin gene, exon 2)					
		and flanking intronic						
1090	AJ224112	sequences	9e-06	<none></none>	<none></none>	<none></none>		
}		Homo sapiens DNA						
1,001	12000212	for repeat sequence		ł	translation initiation factor IF-2	-		
1091	AB000565	Alu	9e-06	72879	Escherichia coli	5.1		
1		H.sapiens flow-sorted						
1 :		chromosome 6		,				
1092	770005	HindIII fragment,			(M65164) 51C surface protein	1		
1092	Z78985	SC6pA20B4	9e-06	159975	[Paramecium tetraurelia]	4.8		
1 1			٠.		50S RIBOSOMAL PROTEIN			
1093	701677	Thermotoga maritima			L2 maritima >gi 437926			
1093	Z21677	DNA for spc operon	9e-06	585879	(Z21677) ribosomal protein L2	7e-14		
		Drosophila hydei			DYNEIN BETA CHAIN,	1		
		Dhc7 (Threads)			CILIARY sea urchin			
1094	AF031494	mRNA, complete cds	0- 06	700277	(Anthocidaris crassispina) chain	[
1024	A1 031494	Homo sapiens	9e-06	729377	[Anthocidaris crassispina]	4e-18		
		placental protein						
1 1		17a1 (PP17) mRNA,						
1095	AF051315	complete cds	4e-06	<none></none>	NONE			
		Homo sapiens	46-00	CNONES	<none></none>	<none></none>		
	-	(subclone 2_f4 from	1					
		BAC H107) DNA			(AE000952) ISA 1214-6,			
1096		sequence	4e-06	2648304	putative transposase	6.2		
		H.sapiens mRNA for			pamerre transposuse	0.2		
	Į.	skeletal muscle-	ľ		(AB016726) calpain	Ī		
1097	X85030	specific calpain	4e-06	4239857	[Schistosoma japonicum]	0.006		
		Human polymorphic			, and a second position of the second positio	0.000		
		arylamine N-				i		
1098	M75162	acetyltransferase	3e-06	<none></none>	<none></none>	<none></none>		
		Rattus norvegicus						
- 1	II	mRNA for CDP-	ĺ			ŀ		
1		diacylglycerol	ł	i		ľ		
,,,,,,		synthase, complete	ļ		(Z70309) R102.6			
1099		cds	3e-06	3879045	[Caenorhabditis elegans]	7.3		
- 1		H.sapiens flow-sorted	1		MERCURIC REDUCTASE			
		chromosome 6	- 1		(HG(II) REDUCTASE)			
,,,,,		HindIII fragment,	1		>gi 418744 pir \$30168	1		
1100	Z78985 S	SC6pA20B4	3e-06	266529	mercury(II) reductase	6.5		
ł	i.	In-marine : Day	1					
1		lomo sapiens mRNA	· ·		i	l		
- 1		or Nedd8-activating	ļ			ł		
1101		nzyme hUba3,	3.00		(Z79697) F58H10.1	ł		
	72017130 C	omplete cds	3e-06	3877938	[Caenorhabditis elegans]	6.3		

	Nearest	Neighbor (BlastN vs. G	enhank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		Treighbor (Blasar Vs. C	T T	Nearest Neigh	noor (BlastX vs. Non-Redundant F	roteins)	
ID	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE	
_		Homo sapiens				Ť	
1		WSCR4 gene, exons	1		(Z80775) hypothetical protein	-f	
1102	AF041056	3 and 4	3e-06	1568583	Rv0044c	1.9	
		Mouse E(d) beta gene					
	Í	5' flanking region and			(U72497) fatty acid amide		
1103	X00777	exon l	3e-06	1680722	hydrolase [Rattus norvegicus]	0.008	
i		Human mRNA for					
		estrogen responsive		}			
1,,,,,	Davage	finger protein.			(U09825) acid finger protein	1 .	
1104	D21205	complete cds	3e-06	563127	[Homo sapiens]	le-05	
1105	747046	Human cosmid					
1103	Z47046	QLL2C9 from Xq28 Human MHC class III	1e-06	<none></none>	<none></none>	<none></none>	
1106	126261	1			1		
1100	L26261	HLA-RP1 gene.	1e-06	<none></none>	<none></none>	<none></none>	
1107	M13402	Rat 5S RNA gene, clone 5S-2.	1. 06	No.		}	
1107	1/11/3402	H.sapiens gene for	1e-06	<none></none>	<none></none>	<none></none>	
1108	X68793	antithrombin III	1e-06	4NIONIT-	NO. =	1	
1.00	100775	antitui Onioni III	16-00	<none></none>	<none> ZINC FINGER PROTEIN 33A</none>	<none></none>	
					(ZINC FINGER PROTEIN	1	
i		Homo sapiens			KOX31) (KIAA0065)	i .	
		Krueppel family zinc			(HA0946) Kruppel-related.		
1109	AF003540	finger protein	1e-06	2507553	[Homo sapiens]	0.098	
		Homo sapiens	10 00	2301333	[[Holito sapiens]	0.098	
i I		(subclone 10_d2 from					
		P1 H21) DNA			(U58762) T27F7.1 gene product		
1110	L42096	sequence.	1e-06	1330401	[Caenorhabditis elegans]	0.015	
		Human DNA			(Cuerrandonna crogaria)	0.015	
ŀ		sequence from	1			l	
ŀ		cosmid cN116A5,	1				
		between markers					
' I		D22S280 and					
ŀ		D22S86 on	ł				
j		chromosome 22q12	ļ		·		
1111	Z69925	contains EST	9e-07	<none></none>	<none></none>	<none></none>	
	İ		. [•			
1			1		(Z81109) predicted using		
- 1	l,	C agrandaine agra Gal	· 1		Genefinder; similar to	į	
J		S. cerevisiae gene for YmL33.	į		sodium/phosphate transporter;		
l i		r mL33, mitochondrial			cDNA EST yk326f6.3 comes	ĺ	
1			1		from this gene; cDNA EST	İ	
1112		ribosomal proteins of	0- 0-		yk326f6.5 comes from this gene		
1114	D90217	arge subunit	9e-07	3879097	[Caenorhabditis elegans]	7.1	

1.5	Nearest	Neighbor (BlastN vs. C	Senhank)	T N		
SEQ		Treighbor (Blastia vs. C	Jenbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant Pr	oteins)
ID ID	ACCESSION	DESCRIPTION	P VALUE	ACCECCION	DECCE PROPERTY.	l
—	1	T DESCRIPTION	FVALUE	ACCESSION	DESCRIPTION	P VALUE
_	 		 	 	(US8755) coded for by C.	
			I	1	elegans cDNA yk34b1.5; coded	
1	1	İ	l		for by C. elegans cDNA	
1			ì		yk13h10.5; coded for by C.	l
1			[elegans cDNA yk46e8.5; coded	
	1	Sambucus nigra	i		for by C. elegans cDNA	
1		ribosome inactivating	ŀ	Į.	yk46d5.5; coded for by C.	i .
		protein precursor			elegans cDNA yk43c2.5; coded for by C. elegans cDNA	·
1113	AF012899	mRNA, complete cds	9e-07	1330345	yk46e8	0.00
			2001	1330343	ук40е8	2e-29
1	j				(U40945) coded for by C.	
1				}	elegans cDNA yk74b9.3; coded	
1	1		٠.		for by C. elegans cDNA	
1	ļ]			yk74b9.5; similar to repeat of	
I			i		calcium channel alpha subunits;	
		g			similar to tetracycline resistance	
1 i		Homo sapiens full		-	protein; similar to hypothetical	- 1
1	. =	length insert cDNA			protein in HSP30-PMP1 region	l l
1114	AF086562	clone ZE16C03	4e-07	1072210	(SP	3.9
ŀ					(Z46/95) similar to	
					transforming protein etc2;	
					cDNA EST EMBL:D34137	
					comes from this gene; cDNA	j
					EST EMBL:D37172 comes	ľ
i 1		Homo sapiens	I		from this gene; cDNA EST	
		interleukin 9 receptor	j		EMBL:D76266 comes from this	
		IL9R pseudogene,	I		gene; cDNA EST	I
1115		exons 1-9	4e-07	3879983	EMBL:D70493 comes from this	
				3079903	gene; cDNA	3.3
]		į			1
		Human DNA	Į.			- 1
		sequence from				1
.		cosmid L96F8,	ı		,	1
- !		Huntington's Disease	1			1
i		Region, chromosome				
J		4p16.3 contains EST	1			
- 1		and cDNA. > ::	j			1
- 1		emb Z69365 HSL96F			i .	1
- 1		BA Human DNA	i		· 1	
J		sequence from		ļ		
- 1		osmid L96F8,	j		1	1
		Iuntington's Disease			1	1
		Region, chromosome	1			
1116		p16.3 contains EST	4- 02		(AF022889) latent TGF beta	1
	209304 B	nd cDNA.	4e-07	3493176	binding protein [Mus musculus]	3.0

	Nearest	Neighbor (BlastN vs. (Genbank)	Neares Nais	hbor (BlastX vs. Non-Redundant F	V
SEQ				inearest ineig	nbor (Blasta, vs. Non-Redundant P	roteins)
ID	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE
	 	Human mRNA for				1
i	1	KIAA0164 gene,			(AC005936) hypothetical	
1117	D79986	complete cds	4e-07	4038031	protein [Arabidopsis thaliana]	0.30
1		Human mRNA for			(and a second analysis)	0.50
1		KIAA0098 gene,	1			ľ
1118	D43950	partial cds	3e-07	<none></none>	<none></none>	<none></none>
i	ł				(AL032657) predicted using	-
1	i				Genefinder; similar to DnaJ	
1	ł		1 .	į.	domain; Thioredoxin; cDNA	İ
1	j	İ			EST yk433f3.5 comes from this	
			İ		gene; cDNA EST	
1		Arabidopsis thaliana	1		EMBL:D32359 comes from this	. • • •
1		DnaJ homologue			gene; cDNA EST	į į
1		(AU6) mRNA,			EMBL:D34721 comes from this	1
1119	AF037168	complete cds	3e-07	3881075	gene; cDNA EST yk433f3.3 c	3e-09
1	******	H.sapiens mRNA for			(U00043) similar to D.	
1120	X69838	G9a	3e-07	3873414	melanogaster trithorax protein	3e-29
i l		Homo sapiens mRNA			1	
1,,,,	4.00	for KIAA0552	i		(U90880) hypothetical protein	1
1121	AB011124	protein, complete cds	2e-07	2618749	2; predicted using XGrail	2.0
		Human cellular fms				
1122	************	proto-oncogene,			į.	i
1122	K03012	partial cds.	1e-07	<none></none>	<none></none>	<none></none>
		Home DNA		•		
		Homo sapiens DNA, microsatellite and Alu				
1123					!!!! ALU SUBFAMILY SQ	
1123	ABOIO193	repeat region	le-07	728837	WARNING ENTRY	0.095
		Homo sapiens			1	
1124		psihHaA pseudogene	45.00	A101m		
		Homo sapiens FLI1	4e-08	<none></none>	<none></none>	<none></none>
- 1		gene for ERGB			İ	Í
I		transcription fuctor,	f	•	. [i
1		intron 4 and partial	ł	•	IIII ATTI GUIDELLE EL CO	ŀ
1125		eds	4e-08	770026	!!!! ALU SUBFAMILY SP	
		Homo sapiens oggl	46-00	728836	WARNING ENTRY	3.6
1126	_ 1	gene, exons 1-7	4e-08	113660	!!!! ALU CLASS C WARNING	
		Homo sapiens	70-00	113668	ENTRY !!!!	3e-05
ļ		subclone 1_c10 from	1			
- 1		P1 H69) DNA	j		ļ	1
1127		equence	3e-08	4225950	(AI132701)	
一十		-1	30-08	7223930	(AJ132701) centaurin gamma1B (AC004537) similar to tumor	1.8
- 1	lo	Gallus gallus mRNA	1			1
- 1		or high mobility	i	İ	suppressor p331NG1; similar to	ł
1128		roup 1 protein	3e-08	3041955	AF044076 (PID:g2829208)	
		Iomo sapiens FGFR-	50-00	3041855	[Homo sapiens]	3e-31
1129		gene	le-08	NONE	· NONE	,,,,,,,
			.0-00	<none></none>	<none></none>	<none></none>

	Near	est Neighbor (BlastN	vs. Genbank)	Nearest Na	ighbor (Pleasy	
SE	Q			. ibarcst ive	ighbor (BlastX vs. Non-Redundar	t Proteins)
ED	ACCESSI	ON DESCRIPTIO	N P VALU	E ACCESSION	DESCRIPTION	PVALUE
						II VALUE
- 1		Mesocricetus aura	ıtus			
- 1	i	serum amyloid P		1		
1112/		component gene,	- 1	İ		
1130	L22024	complete cds.	le-08	<none></none>	<none></none>	3/03-5
ĺ	1	Sambusus air			·	<none></none>
		Sambucus nigra	. (1		
1	1	ribosome inactivat	ing	ì		[
1131	AF012899	protein precursor		· I		
1	A 01289	mRNA. complete	cds 1e-08	<none></none>	<none></none>	<none></none>
ł	1	Human mRNA for	1	<u> </u>		SINONES
- 1	i	phospholipase C >		<i>'</i>	1	
1	ſ	gb M37238 HUMP		1		- 1
- 1	ı	C Human	L	1		
- 1	l	phospholipase C		1		
1132	X14034	mRNA, complete c		İ		
	111.057	H.sapiens CpG DN	ds. 1e-08	<none></none>	<none></none>	<none></none>
	ĺ	clone 152b10,	^· ·			1
1 1		forward read	1		1	1
1133	Z59381	cpg152b10.ft1a.	1	•	1	1 1
		Homo sapiens	le-08	<none></none>	<none></none>	<none></none>
1 1		(subclone 2_h3 from	. 1	l		
		P1 H43) DNA	` [1
1134	L81839	sequence	1e-08	Mone	1	1 1
			16-08	<none></none>	<none></none>	<none></none>
1 1					HYPOTHETICAL PROTEIN	
					MJ1207 Methanococcus	1 1
1 1		i		l	jannaschii >gi 1591837	1 1
1 1		Human GLA gene fo			(U67562) protease synthase and	
1 1		alpha-D-galactosidas	e		sporulation negative regulator	1 1
1135	X14448	A (EC 3.2.1.22)	le-08	2224407	Pail, putative [Methanococcus	1 1
		Human DNA	10-00	3334427	jannaschii]	9.1
		sequence from clone	1 . 1	· .		
		799F15 on] !			
		chromosome Xq25,	1		1	1 1
	l	complete sequence	1 . 1		(150270)	1 1
1136	AL023774	[Homo sapiens]	1e-08	1354935	(U58330) probable copper-]]
1		H.sapiens DNA	 	1334933	transporting atpase	1.2
- 1		repetitive	i i			
		Subtelomeric-like	, i	İ	hungahari 1 gorr	1
1137	X64639	sequence (522 bp)	1e-08	77356	hypothetical 70K protein -	
	[1	Human HuD gene,			eggplant mosaic virus	0.098
1138		UTR	5e-09		(AF070530) unknown [Homo	1
				3307000	sapiens]	9.5

	Neares	t Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant P	
SEC				110410011101	(BlastA vs. :40n-Redundant P	roteins)
ID	ACCESSIO		P VALUE	ACCESSION	DESCRIPTION	P VALU
 		Human DNA				1. 120
ĺ	1	sequence from				
ļ	j	cosmid E86D10 on	i	1	1	
İ	ł	chromosome 22.	1		i	
1	1	contains ESTs,	J			
1139	700.00	exontrap, complete			!!!! ALU SUBFAMILY J	i
1139	Z82181	sequence	5c-09	728831	WARNING ENTRY	8.4
		Mus musculus mRNA	4			
		for translation	1	1		l
1140	AJ006587	initiation factor eIF2	İ		(U22376) alternatively spliced	
1140	AJ000387	gamma X	5e-09	1872200	product using exon 13A	0.64
1141	V11100	H.sapiens WNT8B		• •	(AF045646) contains similarity	-
1141	Y11108	gene	4e-09	2854198	to collagens	4.0
	1	Transmission				
	i	Treponema pallidum section 39 of 87 of	i .		1	
1142	AE001223				CELL DIVISION PROTEIN	
	112001223	the complete genome	4e-09	3334189	FTSY HOMOLOG	1.5
					G1	
					fibroblast growth factor receptor	
1					Al precursor - African clawed	
- 1		Human cosmid			frog >gi 214894 (M55163)	
1143	Z47046	QLL2C9 from Xq28	4e-09	104045	fibroblast growth factor receptor	
		Homo sapiens	46-09	104045	[Xenopus laevis]	1.3
1		genomic DNA, 21q	j			
		region, clone:	1		IIII ALVIGI AGG A WARREN	
144	AG000746	T171Bm40	4e-09	113666	!!!! ALU CLASS A WARNING	
				113000	ENTRY !!!!	0.33
					arginine rich domain, possesses	
	ļ		1		weak similarity with the RNA	
	•]		binding domains from RNA	
		}	1		splicing factor U2AF 65 KD	
- 1	. 1		1		subunit; cDNA EST	ſ
	1	1	· j		EMBL:D64658 comes from this	
ı	. [j	i		gene; cDNA EST	
- 1	ŀ	ľ	1		EMBL:D66829 comes f	
J	j	[I		>gi 3878699 gn1 PID e1351700	
	ĺ	·	- 1		possesses weak similarity with	ı
- }	1	1	J		the RNA binding domains from	ı
1	İ		í		RNA splicing factor U2AF 65	i
- 1	1_		. [KD subunit; cDNA EST	1
		Iuman arginine-rich	j		EMBL:D64658 comes from this	į
15		uclear protein			gene; cDNA EST	
45		RNA, complete cds.	4e-09	3875371	EMBL:D66829 comes f	3e-06
		enopus laevis XL-				-55 55
		NCENP (XL-	1	. [ŀ	ł
46		NCENP) mRNA,			ENDO-1,4-BETA-XYLANASE	ŀ
	U7JU94 C	omplete cds	2e-09	2494337	PRECURSOR sp.]	4.9

	Nearest	Neighbor (BlastN vs. (Genbank)	Negreet Maia	hhor /PlastV N	
SEQ		73. (ivearest iveig	hbor (BlastX vs. Non-Redundant F	roteins)
, ,	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DEGGRA	
			T T VALUE	ACCESSION	DESCRIPTION IUDP-	P VALUE
 		+	 		_1	
		1	1	i	GLUCOSE:GLYCOPROTEIN	
1 1		Drosophila		1	GLUCOSYLTRANSFERASE	
1 1		melanogaster UDP-			PRECURSOR (DUGT)	į p
		glucose:glycoprotein	l		glucosyltransferase - fruit fly (Drosophila sp.)	ļ
		glucosyltransferase			glucosyltransferase precursor	
1147	U20554	mRNA, complete cds.	2e-09	2499087	[Drosophila melanogaster]	4- 24
				2133007	[Diosophila meranogaster]	4e-24
1 1		H.sapiens CpG DNA,				
		clone 91c9, forward			1	1 1
1148	Z56162	read cpg91c9.ft1a.	le-09	··· <none></none>	<none></none>	<none></none>
		Mus musculus				4.01.12
	. =	Pontin52 mRNA,			(U25739) YSPL-1 form 1 [Mus	1
1149	AF100694	complete cds	1e-09	1002424	musculus]	8.9
		U				
1150	M85276	Homo sapiens NKG5			(AF016447) No definition line	1 1
1130	W103270	gene, complete cds. Human	le-09	2315436	found [Caenorhabditis elegans]	8.3
1 1		dihydroorotate				1
		dehydrogenase			LAROUAGO NACE ON THE	1 1
1151	M94065	mRNA, 3' end.	1e-09	3892656	(AB014464) MGC-24v [Mus	
		Homo sapiens	10-07	3092030	musculus]	6.2
		genomic CAG repeat	ľ			1
		element, clone	ı			
1152		60o2(250)	5e-10	<none></none>	<none></none>	<none></none>
1 1		Human DNA			THE THE PARTY OF T	KNONES
		sequence from				
1 1	1	cosmid E86D10 on			l j	
		chromosome 22.	I]	
]]		contains ESTs,			1	1
1153	~~~ 1	exontrap, complete	1		!!!! ALU SUBFAMILY J	
1133		Homo sapiens mRNA	5e-10	728831	WARNING ENTRY	7.9
		or putative				
1154 A		nethyltransferase	50.10	11265	!!!! ALU CLASS B WARNING	
		Homo sapiens RET	5e-10	113667	ENTRY !!!!	0.15
		inger protein-like I	ŀ			- 1
		ntisense transcript,	1		IIII ALII CIIDEANE V coc	
1155 A		artial	5e-10	728834	!!!! ALU SUBFAMILY SB2 WARNING ENTRY	0.001
				140034	WARMING ENTRY	0.006
		łomo sapiens	I			<u> </u>
		ilencer of death	1			1
		omains (SODD)	j		(AF111116) silencer of death	
1156 A		RNA, complete cds	5e-10		domains [Homo sapiens]	2e-08
1		lomo sapiens mRNA			- Carrier Supremoj	20-00
		or hypothetical	1		· .	
1157 2	Z97017 p	rotein	4e-10	<none></none>	<none></none>	<none></none>

-	iveares	st Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant	Proteins
SEC	=				- Tour Road III and III	/ (cicilis)
ID	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	D
		Homo sapiens type I	1			P VALUE
		integral membrane		1		-
1158	8 AF001298	Thi otom	4e-10	<none></none>	<none></none>	NONE
		H.sapiens mRNA for			(U34384) CheW [Borrelia	<none></none>
1159	Y11395	p40	2e-10	1000340	burgdorferi	2.4
		Human non-coding	1			
	ŀ	sequence upstream	1	l		Ĭ
1160	U41096	from DOC-2 gene on chromosome 5	1		!!!! ALU SUBFAMILY SQ	1
1.00	041090	chromosome 5	2e-10	728837	WARNING ENTRY	0.28
	1	Sambucus nigra]			
	I	ribosome inactivating	ĺ	i .	1	
		protein precursor	1	<i>"</i>	}	
1161	AF012899	mRNA, complete cds	6e-11	«NONTE»		
		S.cerevisiae	00-11	<none></none>	<none></none>	<none></none>
	1	chromosome II	,		1	1
	1	reading frame ORF				
1162	Z36111	YBR242w	6e-11	2213560	(Z97052) hypothetical protein	2 25
					(Z80220) Similarity to yeast	3e-27
					protein TREMBL ID E246895)	.1
					cDNA EST EMBL:T00018	
					comes from this gene; cDNA	1 1
		1	I	•	EST EMBL:C13908 comes	}
		Schizosaccharomyces	ı		from this gene; cDNA EST	1
		pombe mRNA, partial	ĺ		EMBL:C11656 comes from this	1 1
1163	D89174	cds, clone: SY 1004	60.11	2.552.55	gene; cDNA EST yk234a5.3	1 1
	20,1,4	Human DNA	6e-11	3879758	comes from this ge	4e-30
- 1		sequence from	1			
		cosmid A1 on	i		ļ	1 1
I		chromosome 6	- 1		1	ĺ
		contains ESTs.			1	1
		HERV like retroviral	1			
164	Z95437	sequence	5e-11	<none></none>	<none></none>	<none></none>
- 1	1	G			3.0.1.2	CHOINES
- 1		Sambucus nigra	j			
		ribosome inactivating	1		,	1
165		protein precursor	_ [(AF106581) contains similarity	
		mRNA, complete cds Human UbA52 gene	5e-11	3886065	to C4-type zinc fingers	4.9
		coding for ubiquitin-	- 1			
ŀ		22 amino acid fusion	j			- 1
66		protein	70.11	NONE		1
		Homo sapiens full	2e-11	<none></none>	<none></none>	<none></none>
		ength insert cDNA	1			. 7
67		lone ZD40G12	2e-11	2134780	apoptosis inhibitor IAP homolog	
				2134/80	- human	3.8

[Nearest	Neighbor (BlastN vs. C	enbank)	Nagross Naist	hb-/DL W D I I D	
SEQ		1	T. C.I.Dallk,	incarest Neigh	hbor (BlastX vs. Non-Redundant Pr	oteins)
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
<u> </u>						T T
1		Homo sapiens mRNA			P53-BINDING PROTEIN	
l		for KIAA0771	ļ	ľ	53BP2 Bbp/53BP2 [Homo	ļ
1168	AB018314	protein, partial cds	2e-11	3024343	sapiens!	2e-11
i		S.cerevisiae			(AC004537) similar to tumor	20-11
1		chromosome XV			suppressor p33ING1; similar to	
		reading frame ORF		4	AF044076 (PID:g2829208)	
1169	Z74972	YOR064c	2e-11	3041855	[Homo sapiens]	2c-40
		Human DNA			(cromo suprens)	26-40
	ł	sequence from				[
1 1		cosmid E86D10 on		i		
1		chromosome 22.	'		1	
		contains ESTs,				
		exontrap, complete	· •	•		
1170	Z82181	sequence	7e-12	<none></none>	<none></none>	-210275
		H. sapiens red cell		THORES	NONE>	<none></none>
		anion exchanger			· ·	
		(EPB3, AE1, Band 3)	į		hypothetical protein - human	
1171	X77738	gene, 3' region	7e-12	2135416	>gi 288145	00.5
		gone, o region	70 12	2133410	7g1/200143	0.012
		medium-chain acyl-			1	
1 1		CoA dehydrogenase			i i	
1 1		{exon 10, intron 10]			[
1 1		[human, Genomic,	1		!!!! ALU CLASS A WARNING	ŀ
1172	S61977	1407 nt]	6e-12	113666	ENTRY !!!!	
		M.musculus DNA for		113000	(X83413) U88 [Human	0.100
1173		HC1 locus	6e-12	854065		200
		protein S=activated		034003	herpesvirus 6]	2e-06
		protein C cofactor	ŀ			j
1		[rats, liver, mRNA,	ŀ		(AF009243) proline-rich Gla	i
1174		3315 nt]	6e-12	2338292		20.10
		Bovine OXT gene for		2330232	protein 2 [Homo sapiens]	3e-10
		oxytocin, 5'	i		(L77967) small proline-rich	
1175		noncoding region	2e-12	1296429		, I
				1270427	protein with paired repeat	4.1
	42	H.sapiens CpG DNA,	j			- 1
J		lone 10h10, reverse	i		(A F030154) pVII (basiles	
1176		ead cpg 10h 10.rt la .	2e-12	2935221	(AF030154) pVII [bovine	
				2933221	adenovirus type 3]	2.8
1	ĺ,	H.sapiens CpG DNA,	f			
- 1	- 1	lone 10h10, reverse	. [(AF027440)	ľ
1177	1	ead cpg10h10.rt1a.	2e-12	2700450	(AF037440) putative 26 kDa	
	220214	cad operonivatia.	-20-12	2708659	protein [Edwardsiella ictaluri]	2.8
	[x	A.musculus h2-			BETA SCRUIN >gi 1015535	I
1178	ľ	alponin cDNA	20.12		(Z47541) beta scruin [Limulus	[
	217545	aipointi CDNA	2e-12	2497945	polyphemus]	2e-0-∔

	Neares	t Neighbor (BlastN vs.	Genbank)	Nearost Nai	abbas (Plast V as No. D.)	-
SEC				regrest Mer	ghbor (BlastX vs. Non-Redundant	Proteins)
ID	-		P VALUE	ACCESSION	DESCRIPTION	P VALUE
		erythropoietin				1. 11.201
		receptor [human,				
1		placental, Genomic,	į		!!!! ALU SUBFAMILY SC	į
1179	S45332	8647 nt]	7e-13	728835	WARNING ENTRY	0.074
- 1					W. Edward Elviki	0.074
1		Sambucus nigra	1	Ì		
	1	ribosome inactivating	1	}	i i	
1		protein precursor	i		1	1
1180	AF012899	mRNA, complete cds	2e-13	<none></none>	<none></none>	<none></none>
						TROITES
	1	Sambucus nigra				1
	i	ribosome inactivating				·
1		protein precursor			}	
1181	AF012899	mRNA, complete cds	2e-13	<none></none>	<none></none>	<none></none>
	1	Harris G G DVI				1
	ł	H.sapiens CpG DNA,				1
1182	Z59509	clone 15a1, reverse			(AL023634) hypothetical	
1102	239309	read cpg15a1.rt1a . Human CYP11B2	2e-13	3150251	protein	0.66
1183	D10170	gene for steroid 18-			!!!! ALU SUBFAMILY SQ	1
1103	D10170	hydroxylase Human MHC class I	2e-13	728837	WARNING ENTRY	3e-05
		molecule (MICB)	·		LINE-I REVERSE	
1184	U65416	gene, complete cds			TRANSCRIPTASE	j l
	005410	Mus musculus	2e-13	126295	HOMOLOG	бе-11
		IHABP gene,				
1185	AJ006031	promoter	8e-14	0120000	hypothetical protein YPL186c -	
		Human gamma-	0¢-14	2132223	yeast	1.1
1 1		sarcoglycan mRNA,	- 1		(U34976) gamma-sarcoglycan	1 1
1186		complete cds	8e-14	1054903	[Homo sapiens] >gi 4239660	
				1034903	sapiens] ACYL-COA	0.034
	1	}			DEHYDROGENASE, VERY-	
i i			i		LONG-CHAIN SPECIFIC	
1	ľ	Rat mRNA for very-	I		(VLCAD) >gi 2388724	
	ļ:	ong-chain Acyl-CoA	1		(AF017176) very-long-chain	·
	ļ	iehydrogenase,			acyl-CoA dehydrogenase [Mus	
1187	D30647	complete cds	8e-14	3183512	musculus)	8e-23
ŀ						06-23
ı		I.sapiens CpG DNA.	J			ı
		lone 7g4, forward	1			
1188	Z63247 r	ead cpg7g4.fla.	6e-14	86285	histone H1.01 - chicken	6.8
J	را	Seller estimation				
- 1		Gallus gallus zinc	1			l
1189		inger protein (Fzf-1)	1		zinc finger protein - chicken	J
		RNA, complete cds.	3e-14	2134436	(fragment)	4e-10
ĺ		frican green	1			
1190		onkey origin of	20.14	.,,,,,_		í
		pheation	2e-14	<none></none>	<none></none>	<none></none>

		Neare	t Neighbor (DlassN (7a-b13	Т		
	SEQ	1,404165	st Neighbor (BlastN vs. (Jenbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant I	Proteins)
	ID ID	ACCESSIO		P VALUE		DESCRIPTION	P VALUE
ı		 	Mus musculus				1 VALUE
	1191	AF100694	Pontin52 mRNA,			(AF119040) NL0D	
ł	1191	AF100694	complete cds	2e-14	4235641	[Lycopersicon esculentum]	0.65
	1192	AF012899	Sambucus nigra ribosome inactivating protein precursor mRNA. complete cds	2 e -14	3043728	(AB011174) KIAA0602 protein	0.28
	1193	AJ005866	Homo sapiens mRNA for putative Sqv-7- like protein, partial	2e-14	·· 4008517	(AJ005866) Sqv-7-like protein [Homo sapiens]	0.004
	1194	U32709	Haemophilus influenzae Rd section 24 of 163 of the complete genome	 2e-14	3861056	(AJ235272) POLYRIBONUCLEOTIDE NUCLEOTIDYLTRANSFERA SE (pnp) [Rickettsia prowazekii]	
	195	AF073485	Homo sapiens MHC class I-related protein MR I precursor (MR I) gene, partial cds Homo sapiens clone	8 c -15	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	6e-28
1	196	AF052135	23625 mRNA sequence	8e-15	4098124	(U73522) AMSH [Homo sapiens]	8e-14
1	197	AF100694	Mus musculus Pontin52 mRNA, complete cds	3e-15	<none></none>	<none></none>	<none></none>
1	198		Sambucus nigra ribosome inactivating protein precursor mRNA, complete cds	3e-15	113671	!!!! ALU CLASS F WARNING ENTRY !!!!	1.7
11	00		S.cerevisiae chromosome XV eading frame ORF			(Z46381) similar to lipoic acid synthase; cDNA EST yk283b6.3 comes from this gene; cDNA EST yk283b6.5 comes from this gene; cDNA EST yk472f5.3 comes from this gene; cDNA EST yk472f5.5 comes from this	
-11	77	Z75104	OR196c	3e-15	3878570	gene: cDNA EST yk476e7.3	le-15

		t Neighbor (BlastN vs.	Genbank)	Nearest Neig	hbor (BlastX vs. Non-Redundant	Proteins)
SE	`]		1			- Totalis,
ID	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
				T	(U42833) coded for by C.	PVALUE
					elegans cDNA cm16f6; coded	
1				ĺ	for by C. elegans cDNA	
- }			1		CEESU63F; similar to S.]
1		J			cerevisiae SOF1 protein	j
1200	X70052	S.cerevisiae sof1		İ	(SP:P33750) [Caenorhabditis	1
120	7 A70032	gene	3e-15	1125754	elegans]	3e-29
1		Sambucus nigra	i i	Ì		
1		ribosome inactivating	,		1	1
1		protein precursor	'		1	1
1201	AF012899	mRNA, complete cds	2e-15	· · <none></none>	NONE	1
			1 20 13	4101123	<none></none>	<none></none>
1	ł	Gorilla gorilla gamma	1			1
	ł	1 and gamma-2				1
1,000		globin genes,	1		hypothetical protein 2 - human	1
1202	M92295	complete cds.	le-15	284078	>gi 182220	7.4
1		Homo sapiens RNA				
		polymerase II	1			
]	elongation factor SIII,			1	
	Ī	p15 subunit mRNA,	1 1		1	1 '
		complete cds. > ::	ľ	•		1 1
1		gb AR022286 AR022	ĺ		İ	1
	,	286 Sequence 7 from				1 1
1203	L34587	patent US 5792634	9e-16	<none></none>	<none></none>	<none></none>
						CHOINES
		Xenopus laevis				1
1204	D83649	mRNA for xSox7			(D83649) xSox7 protein	
1204	D83049	protein, complete cds Homo sapiens PAC	8e-16	2447043	[Xenopus laevis]	4e-06
		clone DJ1152D16	I			
		from Xq23; complete	1		1 '	1
		sequence [Homo				. [
1205		sapiens]	3e-16	<none></none>	NONE	270275
		Human UMP		VIVOINES	<none></none>	<none></none>
		synthase mRNA.	•		!!!! ALU CLASS B WARNING	
1206		complete cds.	3e-16	113667	ENTRY !!!!	0.65
		Human Alu family				0.05
1207		interspersed repeat;		•	!!!! ALU SUBFAMILY SP	1
1207	J00083	clone BLUR11.	3e-16	728836	WARNING ENTRY	4e-06
ı	1,	Mus musculus m-	ł			
		Numb (m-nb) mRNA.	1	i		1
1208		complete cds	le-16	NONE		j
		pioto cus	16-10	<none></none>	<none></none>	<none></none>

	Negree	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC		TACIBIDOL LD MASHA AS. O	I I	Nearest Neigh	npor (BlastX vs. Non-Redundant Pi	roteins)	
SEQ							
	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
 		Human SWI/SNF	ļ				
	İ	complex 60 KDa	-	Į.	İ	1	
ĺ	ľ	subunit (BAF60c)		<u> </u>		1	
1209	U66619		, ,,		(U66619) SWI/SNF complex 60		
1209	000019	mRNA, complete cds Drosophila	1e-16	1549247	KDa subunit [Homo sapiens]	0.003	
	ı	melanogaster Rga and			+		
		Atu genes, complete			(1775 ((7)) (7)		
1210	U75467	cds	le-16	1650500	(U75467) Atu [Drosophila		
1210	073407	cus	16-10	1658503	melanogaster]	5e-32	
		Human alternative					
		splicing factor			1		
1211	M72709	mRNA, complete cds.	3e-17	<none></none>	ANIONTE:		
		Human ferritin H	<u> </u>	CHOILES	<none></none>	<none></none>	
	!	(FTHL13)			İ		
1212	U26556	pseudogene.	3e-17	<none></none>	<none></none>	<none></none>	
		Human gene for 2-		CHOILE	CHOILE	KINOINES	
		oxoglutarate			(AF003386) F59E12.9 gene		
	•	dehydrogenase,			product [Caenorhabditis		
1213	D32064	complete cds	3e-17	2088843	elegans]	0.12	
				2000045	APAG PROTEIN	0.12	
		Human (Papua New			>gi 72927 pir BVECAG apaG		
		Guinean)			protein - Escherichia coli		
i		Mitochondrial DNA			>gi 40918 (X04711) URF		
ł		control region,	i		hypothetical protein		
1214	M76364	sequence 131.	3e-17	114009	[Escherichia coli]	0.006	
ł		Homo sapiens				-	
ı		genomic sequence	Ī		1		
		from subtelomeric	į				
		region of	ı		(U78948) MADS-box protein 2		
1215	AF017466	chromosome 4q	le-17	3947985	[Malus domestica]	4.1	
	ĺ	.,	į				
		Homo sapiens	Ī				
1214		54TMp (54tm)	· , <u>, </u> [(AF004876) 54TMp [Homo	!	
1216		mRNA, complete cds	le-17	4101574	sapiens]	0.006	
- 1		Pontin52 mRNA.	ľ				
1217		complete cds	1	NO. ==			
	AT 100094	ompiete cas	9e-18	<none></none>	<none></none>	<none></none>	
	Į,	Rattus norvegicus Na-	- 1		(AL033545) putative glycine-	į	
1218		₹-2Cl cotransporter	40.10	3803703	rich protein [Arabidopsis	2.22	
		domo sapiens	4e-18	3892703	thaliana) (AF023910) DNA	0.30	
		PENIIB mRNA.	1			Ī	
219	1	omplete cds	4e-18	2642402	topoisomerase I [Physarum		
		I.sapiens IRLB gene	46-10	2642493	polycephalum] irlB protein - human (fragment)	0.083	
220		exon1-3)	4e-18	106927		2011	
		C.COII 1-3/	46-10	106837	>gi 33969	2e-11	

<u> </u>	Near	est Neighbor (BlastN	vs. Genbank)	Nearest N	leighbor (BlastX vs. Non-Redundan	D
SE	Q				Vs. Non-Redundan	Proteins)
IL	ACCESS	ON DESCRIPTIO	N P VALI	IE ACCESSIO		
		Human mRNA fo		JE ACCESSIC	DESCRIPTION	P VALI
		KIAA0385 gene,				
122	I AB0023	33 complete cds	4e-18	77777	(AF060181) zinc finger prote	n
122	2 X98485	P.vivax PV14 ger	ie le-18	3220370	[Homo sapiens]	6e-25
		H.sapiens flow-so	rted	<none></none>	<none></none>	<none< td=""></none<>
	1	chromosome 6		į		
	1	HindIII fragment,	ł	1	(AD012023) 05-5 1-	j
122	Z79057	SC6pA21E8	le-18	2981631	(AB012223) ORF2 [Canis	
	1			2901031	familiaris]	0.001
	ł	Homo sapiens (clo	ne	1	nucleolar 100K polymyositis-	
	1	JH4B1) PM-scl	1	1	scleroderma protein - human	}
	1	autoantigen mRNA	۸,	1.	>gi 35555 (X66113) PM/Scl	
1224	L01457	complete cds.	le-18	346287	100kD nucleolar protein [Hom	
	1	Dog nonerythroid		3.70207	sapiens] (AB017037) nonstructural	100.0
1000	1	beta-spectrin mRN	Α,		protein precursor [Himetobi P	1
1225	L02897	3' end.	4e-19	3493358	virus]	1
					1411431	0.12
	ſ	Homo sapiens mRN	NA			1
1226	4.00.00.00	for APCL protein,			(AB012162) APCL protein	1
220	AB012162	complete cds	4e-19	3894265	[Homo sapiens]	0.000
		11				0.002
		Homo sapiens mRN	IA	ŀ	1	1
227	AB011093	for KIAA0521		1	(AB011093) KIAA0521 proteir	.1
=-/	AB011093	protein, partial cds X.laevis AB21	4e-19	3043566	[Homo sapiens]	9e-09
I		mRNA for RPD3	ſ	·		70-03
228	X78454	homologue		j	HISTONE DEACETYLASE	ŀ
	1170434	nomotogue	4e-19	3023945	(HD) thaliana]	5e-34
- 1		Human endogenous	1	1		
Į		retrovirus H DI	1	į	}	1
		leader		1		
- 1		region/integrase-	1	1	1	
- 1		derived ORFI,	1	i		
		ORF2, and putative	}		177.42.0	
- 1		envelope protein			(Z14310) tripartite fusion	
29	U88895	mRNA, complete cds	2e-19	50077	transcript PLA2L [Human	i
		Human tyrosine	1	59977	endogenous retrovirus]	le-04
		kinase TXK (txk)			IIII AT II SIMBOO	
30	U34377	gene, exon 13.	le-19	728831	!!!! ALU SUBFAMILY J	
- 1				740031	WARNING ENTRY	3e-05
1			 		(Z99167) putative peroxisomal organisation and biogenesis	
		M.musculus rab3A	!		protein [Schizosaccharomyces	j
31	X72966	gene	le-19	2408076	protein (Schizosaccharomyces pombe)	
1				2.03070	[POINTOE]	2e-09
1		Homo sapiens	l l		1	
- 1		mRNA, chromosome	1		1	1
		specific transcript			1	1
2 /		KIAA0484	1		j · i	I

<u> </u>	Neares	st Neighbor (BlastN vs. (Genbank)	Negree Mail	the Oliver	
SEQ	1		T	Nearest Neis	ghbor (BlastX vs. Non-Redundant Pr	oteins)
ID	ACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALU
					(AB001535) similar to	I VALU
		Human gene for Zn-alpha2-glycoprotein,		·	C.elegans hypothetical protein CET01H8.1,CEC05C12.3,CEF5	
1233	D14034	complete cds H.sapiens HOK-2	2e-20	<u> </u>	4D1.5. similar to trp and trp-like proteins [Homo sapiens]	1.07
1234	X82126	gene, exon 2	2e-20	2137269	DNA-binding protein - mouse >gi 437444	le-07
1235	AF093684	Luciferase reporter vector pXP2 *SA, complete sequence	5e-21	2773363	(AF041382) microtubule	le-19
236	J05272	Human IMP dehydrogenase type I mRNA complete cds.	5e-21	124417	binding protein D-CLIP-190 INOSINE-5'- MONOPHOSPHATE DEHYDROGENASE 1 (IMP DEHYDROGENASE 1) (IMPDH-I) (IMPD 1) I - human	5 .5
237	D0404-	Human (lambda) DNA for immunoglobulin light chain	5e-21	3878261	(Z75712) Similarity to S. Pombe BEM1/BUD5 suppressor; cDNA EST EMBL:Z14470 comes from this gene; cDNA EST yk482d4.3 comes from this gene; cDNA EST yk482d4.5 comes from this gene [Caenorhabditis elegans]	2e-04 6e-46
8 2	c is M	d.sapiens hromosome 22 CpG sland DNA genomic dsel fragment, clone 02f3, forward read 02f3.f	2e-21	, , , ,	(AF024614) ADAM 10 [Caenorhabditis elegans] Zincbinding metalloprotease domain; cDNA EST CEMSA42F comes from this gene; cDNA EST yk218f3.3 comes from this gene; cDNA EST yk443d9.3 comes from this gene; cDNA EST yk443d9.5 comes from this	
	13,	· J . I	∠e-21	2739037 lo	ene; cDNA	2.6

-	Ne	earest N	leighbor (BlastN v	s. Genbank)		Nagrace 3	Visit III	
SE	-Q	ı				ivearest r	Veighbor (BlastX vs. Non-Redundan	Proteins)
	D ACCES	SION	DESCRIPTION	V P VAL	UE	ACCESSIO	ON DESCRIPTION	
							מתווופרטו ובחווווב (כפכסטבובי)	P VAI
1	- 1	- 1		ı			B; cDNA EST vk450d8.5 com	Tes
1	1	- 1		1			from this gene: cDNA EST	
1	· I	ł		- 1	I		yk249a6.5 comes from this	1
1	1	- 1			- 1		gene; cDNA EST yk219a2.5	1
l	ł			i	- [comes from this gene; cDNA	- 1
1	1	- 1		- 1	- 1		EST yk355e4.5 comes from th	is
ł	1	- 1		·	- 1		gene; cDNA EST yk224f4.5	
ľ	I	- 1		1	- 1		comes fr	
	1	- 1		1			>gi 3924881 gnl PID e1354569	.
	j	- 1		1	- [•	from this gene; cDNA EST	1
	1	- 1		1	- "		yk249a6.5 comes from this] .
	1	- 1			- 1		gene; cDNA EST yk219a2.5	İ
	í		us musculus	j	- 1		comes from this gene; cDNA	1
1020	l	Po	ntin52 mRNA,	1	1		EST yk355e4.5 comes from this	
1239	AF10069		mplete cds	6e-22	1	3924779	gene; cDNA EST yk224f4.5	1
1240		Hu	man primary Alu		 	3724779	comes from	0.35
1240	U67824		nscript	6e-22	1	728832	!!!! ALU SUBFAMILY SB	
	i	Ho	mo sapiens clone		1-	720032	WARNING ENTRY	5e-07
241	AF07063		86 mRNA	1	1		fatty-acid synthase (EC	
	71.07063	o seq	uence	2e-22	1	98710	2.3.1.85) - Brevibacterium ammoniagenes	
- 1		и	700 cm of 7	1			ammonagenes	2.5
		alah	nan gene for Zn- na2-glycoprotein,	1	1			
242	D14034	com	plete cds		1		(Y17832) pol protein [Human	
Т		Hun	nan lactase	2e-22	<u> </u>	1185939	endogenous retro virus K	0.29
- 1			rizin hydrolase		l .			0.29
243	M61835	ac	Γ) gene, exon 2.	2e-22	Ι.		!!!! ALU SUBFAMILY J	
- 1		T	2, 3,0,12.	20-22	 	728831	WARNING ENTRY	0.006
- [musculus		1		RABPHILIN-3A	
	4		in52 mRNA,				>gi 477100 pir A48097	ı
44	AF100694		lete cds	6e-23	1.	350828	rabphilin-3A - bovine	
- 1		Hom	o sapiens full		<u></u>	330028	>gi 285646 gnl PID d1003285	0.14
45	A E07400 -		n insert cDNA	j		\	(AE0selle)	
~ }	AF074985	YH73		8e-24	31	70548	(AF056116) unknown (Fugu	
			in mRNA for				rubripes]	0.24
16	D14878		n D123,				1]
+	2140/0	compl	ete cds	7e-24	<n< td=""><td>ONE></td><td><none></none></td><td></td></n<>	ONE>	<none></none>	
-		1		T			(U61955) contains multiple	NONE>
-		Huma	n HepG2 3'	1			region of strong similarity to	
1		region	cDNA, clone	j			C2H2-type zinc fingers	- 1
7		hmd3d					(PS:PS00028) [Caenorhabditis	- 1
			<u> </u>	6e-24	130	7345	elegans	- 1

\vdash		st Neighbor (BlastN vs.	Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant F	Proteins)
SE	-				· · · · · · · · · · · · · · · · · · ·	iolens)
II	ACCESSIO		P VALUE	ACCESSION	DESCRIPTION	
		Human DNA		1	DESCRIPTION	P VALU
		sequence from		 		
	1	cosmid L98A6.	1	1		
	ı	Huntington's Disease		4		
1,24	3,0,0	Region, chromosom	е	l	(AF123462) neurexin III [Homo	
124	8 Z69654	4p16.3.	3e-24	4240566	sapiens)	4.5
1	1	Home series - DM	.			1 4.5
]		Homo sapiens mRN. for KIAA0445	^A			1
1249	AB007914		1	ł	(AF095568) amelogenin	ł
	112007714	protein, complete cas	2e-24	3885949	[Paleosuchus palpebrosus]	3.2
	1		1		immunodominant microneme	
ĺ	I	Homo sapiens full	1	•	protein Etp 100 - Eimeria tenella	1
	1	length insert cDNA			>gi 2707733 (AF032905)	
1250	AF088072	clone ZD93D10	2e-24	323091	microneme protein precursor	
		Homo sapiens cAMP		323091	Etmic-1 [Eimeria tenella]	0.34
	I	specific	1 1			
	I	phosphodiesterase 4A	·		İ	
	j	variant pde46			1	
	,	(PDE4A) gene, exons	1		1	
	ľ	2 through 13 and			1	
		alternative splice	1 1		1	
1251	AF069489	exons 3a, 6a, 6b, and 9a			!!!! ALU SUBFAMILY SP	
	111 007 487	Homo sapiens P2X7	2e-24	728836	WARNING ENTRY	le-05
1252	Y12853	gene, exon 4-8	9e-25		!!!! ALU SUBFAMILY J	
		Human 28S	96-23	728831	WARNING ENTRY	le-05
		ribosomal RNA gene,	1			
1253	M27830	complete cds.	8 e -25	<none></none>	NONE	
				- Trong	<none></none>	<none></none>
		Homo sapiens				I
- 1	1	mRNA, chromosome	1		1	
254	4 B007053	l specific transcript				
234	AB007953	KIAA0484	8e-25	<none></none>	<none></none>	NONE>
		H.sapiens CpG DNA,	1			
- 1		clone 195c8, forward	1			1
255	260212	read cpg195c8.ft1a.	8e-25	150154	(M81959) POU domain protein	j
		Mus musculus	96-73	158154	[Drosophila melanogaster]	3.3
	ľ	Pontin52 mRNA,	1	ł		7
256	AF100694	complete cds	7e-25	<none></none>	aNO) TO	
		Mus musculus		- CHOMES	<none> <</none>	NONE>
		Pontin52 mRNA,	I			
257		complete cds	7e-25	<none></none>	<none> <</none>	NONE.
		Iomo sapiens P2X7			3,0,112	NONE>
- 1		ene, exon 1 and oined CDS				
58	Y12851 j		2e-25			

-		st Neighbor (BlastN vs	. Genbank)	Nearest Nei	ighbor (BlastX vs. Non-Redundant	D
SE	Q				- Non-Redundant	Proteins)
Ln	ACCESSIO	DESCRIPTION	P VALUE	4.0000000		
		Mus musculus Tera	_ L VALUE	ACCESSION	DESCRIPTION	P VALU
		(Tera) mRNA,	·	 		
125	59 U64033		9e-26			
		Rattus norvegicus	96-26	<none></none>	<none></none>	<none:< td=""></none:<>
ı	1	Rabin3 mRNA,	İ	i		
126	0 U19181	complete cds.	9e-26	624226	(U19181) Rabin3 [Rattus	1
			76-20	624225	norvegicus] SEL-10 PROTEIN Candida	le-13
	ł	1		i	CDC4 gene (TR:E234056);	
1	1			Í	cDNA EST EMBL:D27699	1
1	1				comes from this gene; cDNA	J
1	i				EST EMBL: D27698 comes	
ſ	Į		1		from this gene; cDNA EST	1 .
1	1	Caenorhabditis			EMBL:D32793 comes from this	. `
1	i	elegans SEL-10 (sel-			gene; cDNA EST	s
		10) mRNA, complete	e		EMBL:D33271 comes from this	. [
126	AF020788	cds	9e-26	3915881	gen	I .
ı	İ				Echi	7e-32
	i	Cricetulus griseus	1 1			
	1	mRNA for	1			1
		Phosphatidylglycerop)		(AB016930)	1
1262	ABOLGOOD	hosphate synthase,	1 1	·	Phosphatidylglycerophosphate	l
1202	AB016930	complete cds	8e-26	4159682	synthase [Cricetulus griseus]	0.045
		Mus musculus	1 1		(Z93385) predicted using	0.045
1263	AF100694	Pontin52 mRNA,	1		Genefinder; Similarity to	
1203	AT 100094	complete cds	3e-26	3878629	B.subtilis GTP-binding protein	2e-10
1264	X91195	H.sapiens SOM172 mRNA	1 1			
		Mus musculus	le-26	<none></none>	<none></none>	<none></none>
	Î	Pontin52 mRNA	1			
1265	AF100694	complete cds				í
	1300)	Human MG21	Ie-26	1360637	(X95995) ENBP1 [Vicia sativa]	3.1
1266	L08237	mRNA. partial cds.	10.26	0.70	(L08237) located at OATL1	
		nadori, partiar cus.	Ie-26	950411	[Homo sapiens]	9e-09
l					(41.032657)	
- 1			1	•	(AL032657) similar to EGF-like	•
]	J		domain; cDNA EST yk299a12.3	į
J		1	j		comes from this gene; cDNA EST EMBL:D35398 comes	1
ı	ł	1	1			- 1
- 1	1]		from this gene; cDNA EST	
- 1		Mus musculus	J		yk331h6.5 comes from this	
	1	Pontin52 mRNA,	1		gene; cDNA EST yk299a12.5	
267		complete cds	9e-27		comes from this gene; cDNA	
		Mus musculus		200.000	EST yk467g8	100.0
J	1.				1	
268		Pontin52 mRNA, complete cds	i i]	HYPOTHETICAL PROTEIN	i

	Neares	t Neighbor (BlastN vs. (Genbank)	Nearest Neig	thbor (BlastX vs. Non-Redundant P	
SE					THE THE THE THE THE THE THE THE THE THE	oteins)
ID	` [N DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE
—	-					
		H sapiens DNA for	1			
1,00		endogenous retroviral			(Y12713) Gag polyprotein [Mus	l
126	9 X89211	like element	8e-27	2065209	musculus]	0.005
1	1	TTomos and				
-	1	Homo sapiens cosmid				
1		clone LUCA15 from			1	Ì
		3p21.3, complete	l	1	1	
1270	U73166	sequence [Homo			!!!! ALU SUBFAMILY J	
12/	0/3100	sapiens	3e-27	728831	WARNING ENTRY	4e-04
		Mouse mRNA for		İ		
1271	D78255	PAP-1. complete cds	2 . 07		(D78255) PAP-1 (Mus	
	270233	Mus musculus	3e-27	1850098	musculus]	2e-10
		Pontin52 mRNA.				
1272	AF100694	complete cds	le-27	2122570	spermatophorin Sp23 - yellow	
		complete eds	16-27	2133579	mealworm molitor	0.39
	· ·	Homo sapiens gene			(Z83318) predicted using	
1		for hippocalcin, exon			Genefinder; cDNA EST	
1273	AB015202	2, 3 and complete cds	Ie-27	3877698	yk369e7.5 comes from this gene	
		Mus musculus	10-27	3877098	[Caenorhabditis elegans]	0.37
		Pontin52 mRNA.			(AF074902) laminin alpha chain	
1274	AF100694	complete cds	le-27	3328188	[Caenorhabditis elegans]	
		H.sapiens gene for		3320130	(Cachornabuttis elegalis)	0.19
		Cu/Zn-superoxide	ľ		!!!! ALU SUBFAMILY J	j
1275	Z29336	dismutase	_le-27	728831	WARNING ENTRY	6e-05
		Mus musculus				-00-03
		Pontin52 mRNA,	I		spermatophorin Sp23 - yellow	
1276	AF100694	complete cds	9e-28	2133579	mealworm molitor)	9.2
		Mus musculus				
1227	A.E.1.00.co.	Pontin52 mRNA,			spermatophorin Sp23 - yellow	i
1277	AF100694	complete cds	9e-28	2133579	mealworm molitor]	0.054
i			1		PUTATIVE PRE-MRNA	
ľ		Wanna and and Day a	I		SPLICING FACTOR ATP-	ſ
1		Homo sapiens mRNA	1		DEPENDENT RNA	1
		for ATP-dependent RNA helicase #46,	Í		HELICASE >gi 2275203	1
1278					(AC002337) RNA helicase	
	112001030	complete cds	4e-28	3913425	isolog [Arabidopsis thaliana]	3e-22
			i		(AC005990) Contains repeated	
l	,		I	,	region with similarity to	
ı		j	- 1		gb U43627 extensin (atExt1)	
- 1	1,	Mus musculus	- 1		gene from Arabidopsis thaliana.	
	1	Pontin52 mRNA,			ESTs gb Z34165 and gb Z18788	
1279		complete cds	3e-28	4056454	come from this gene.	
		7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	30-20	4030434	[Arabidopsis thaliana]	0.066

-		st Neighbor (BlastN v.	s. Genbank)	Nearest Nei	ghbor (BlastX vs. Non-Redundant P	rotairs)
SEC	2				2	roteins)
ID	ACCESSIO	DN DESCRIPTION	P VALUE	ACCESSION		P VALU
-		- 			(AC005990) Contains repeated	
1	1				region with similarity to	
ĺ	ł		1	ł	gb U43627 extensin (atExt1)	
1	ł	Management	ľ		gene from Arabidopsis thaliana.	
1		Mus musculus Pontin52 mRNA,	- I		ESTs gb Z34165 and gb Z18788	
1280	AF100694			1	come from this gene.	1
	A 100094	Mus musculus	3e-28	4056454	[Arabidopsis thaliana]	4e-05
	Í	Pontin52 mRNA.				10 00
1281	AF100694					1
	74 100094	Mus musculus	1e-28	<none></none>	<none></none>	<none></none>
	Į.	Pontin52 mRNA,		}		
1282	AF100694	complete cds	1			
	100094	Mus musculus	le-28	<none></none>	<none></none>	<none></none>
	}	Pontin52 mRNA.				
1283	AF100694		1- 20			
		Mus musculus	le-28	<none></none>	<none></none>	<none></none>
		Pontin52 mRNA,	1			
1284	AF100694	complete cds	le-28		1	
		Mus musculus	16-28	<none></none>	<none></none>	<none></none>
- 1		Pontin52 mRNA,				
1285	AF100694	complete cds	le-28	-NONT		
		Mus musculus	10-26	<none></none>	<none></none>	<none></none>
1		Pontin52 mRNA,				
1286	AF100694	complete cds	1e-28	<none></none>	1	
- 1			1 - 1	- CIVOIVE	<none> PROBABLE INTRON</none>	<none></none>
- 1		Mus musculus	1 1		MATURASE liverwort	
- 1		Pontin52 mRNA,	1	ı	(Marchantia polymorpha)	ŀ
287	AF100694	complete cds	le-28	140505	chloroplast >gi 11663	ا م
					PROBABLE INTRON	3.0
1	:	Mus musculus	1		MATURASE liverwort	ľ
		Pontin52 mRNA,	1		(Marchantia polymorpha)	ľ
288	AF100694	complete cds	le-28	140505	chloroplast >gi 11663	1.8
- 1		Mus musculus			and the second s	1.8
289		Pontin52 mRNA,	1		spermatophorin Sp23 - yellow	
209	AF100694	complete cds	le-28	2133579	mealworm molitor]	0.50
					(AC005990) Contains repeated	0.50
ł	J				region with similarity to	
1	ļ	i			gb U43627 extensin (atExt1)	ı
	1,	Mug musaulus			gene from Arabidopsis thaliana.	
1		Mus musculus	- 1		ESTs gb Z34165 and gb Z18788	1
90		Pontin52 mRNA,	1		come from this gene.	- 1
+	11 100094	complete cds	1e-28	4056454	[Arabidopsis thaliana]	0.087
	1,	Lsapiens CpG DNA.	J			
j		lone 77b3, forward	1	İ	HYPOTHETICAL 29.3 KD	1
91		ead cpg77b3, ft1a.	10.20		PROTEIN pseudotsugata	
		operros.mia.	le-28	2493240	nuclear.polyhedrosis virus]	0.014

<u> </u>	Neare	st Neighbor (BlastN v.	s. Genbank)	Nearest Ne	ighbor (BlastX vs. Non-Redundant)	
SEQ				7.00.051110	Ignoof (BlastA vs. Non-Redundant)	Proteins)
ID	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION		P VALU
	1				DEHYDRIN DHN3	
l	Í	Mus musculus		1	>gi 100035 pir S18139 dehydri	n
l	1	Pontin52 mRNA.	1	1	DHN3 - garden pea >gil20709	1
1292	AF100694		1.00		(X63063) pea dehydrin DHN3	1
		Testificie cus	le-28	118588	[Pisum sativum]	0.010
				ł	(AC005990) Contains repeated	
			J	1	region with similarity to	1
				Į.	gb U43627 extensin (atExt1)	1
		Mus musculus	J		gene from Arabidopsis thaliana.	I
		Pontin52 mRNA,		1	ESTs gb Z34165 and gb Z18788	
1293	AF100694	complete cds	le-28	4056454	come from this gene.	1
- 1	<u></u>		1	+030434	[Arabidopsis thaliana] (AC005990) Contains repeated	0.007
- 1					region with similarity to	
- 1		1			gb U43627 extensin (atExt1)]
			1		gene from Arabidopsis thaliana.	j
		Mus musculus	1		ESTs gb Z34165 and gb Z18788	
1294	4 E1 00 co .	Pontin52 mRNA,	1 1		come from this gene.	
294	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana]	0.000
I		Mus musculus			LAMININ ALPHA-I CHAIN	0.002
295	AF100694	Pontin52 mRNA,	1		PRECURSOR precursor -	
	AL 100094	complete cds	le-28	126363	human	3e-04
1		ĺ	1 1	· ·	(AC005990) Contains repeated	36-04
1			1 1		region with similarity to	
1			1 1		gb U43627 extensin (atExt1)	
- 1		Mus musculus	1 1		gene from Arabidopsis thaliana.	
- 1.		Pontin52 mRNA,	!!!		ESTs gb Z34165 and gb Z18788	
296	AF100694	complete cds	le-28	1056.45.	come from this gene.	
Т			16-28	4056454	[Arabidopsis thaliana]	le-04
- 1	j				(AC005990) Contains repeated	
- [- 1		l		region with similarity to	
- 1	į		- 1		gb U43627 extensin (atExt1)	
- 1		Mus musculus	. 1	•	gene from Arabidopsis thaliana.	l
		Pontin52 mRNA,	I		ESTs gb Z34165 and gb Z18788	
97 A	AF100694	omplete cds	le-28	4056454	come from this gene.	
- 1				.050154	[Arabidopsis thaliana] (AC002131) Strong similarity to	3e-05
-		Aus musculus	1		extensin-like protein gb Z34465	l
98 A		ontin52 mRNA,			from Zea mays. [Arabidopsis	[
'° A	F100694 c	omplete cds	le-28	3157926	thalianal	a. 0.
					(AC005990) Contains repeated	2e-05
1			[ļ	region with similarity to	- [
1			l	ł.	gb U43627 extensin (atExt1)	1
	J.	fue museul.	1	1:	gene from Arabidopsis thaliana.	
		lus musculus	I	ļi	ESTs gb Z34165 and gb Z18788	
9 AI		ontin52 mRNA, omplete cds		10	come from this gene.	J
	.00074 [60	implete cas	le-28		[A = b. 4	1e-05

-		st Neighbor (BlastN)	vs. Genbank)	Nearest Ne	ighbor (BlastX vs. Non-Redundant P	
SE	Q			1	VS. Non-Redundant P	roteins)
II	ACCESSIO	DESCRIPTION	N PVALUE	ACCESSION	The state of the s	P VAL
		Mus musculus			kinetoplast-associated protein -	
l		Pontin52 mRNA.			Trypanosoma cruzi >gil162142	
130	0 AF100694			Í	(M25364) kinetoplast-associated	ı
	12.000	Complete cus	le-28	320919	protein	le-07
	ł		1		(AC005990) Contains repeated	
1	I		ł	1	region with similarity to	1
ı	i				gb U43627 extensin (atExt1)	l
	1	Mus musculus			gene from Arabidopsis thaliana.	
	[Pontin52 mRNA,	1	j	ESTs gb Z34165 and gb Z18788	
1301	AF100694	complete cds	le-28	105615	come from this gene.	
		- Provide Edg	16-28	4056454	[Arabidopsis thaliana]	9e-08
	1	1	}]	(AC005990) Contains repeated	
	1	ľ		[region with similarity to	
	i	1	ı I	İ	gb U43627 extensin (atExt1)	
	l	Mus musculus			gene from Arabidopsis thaliana.	
	1	Pontin52 mRNA,	1		ESTs gb Z34165 and gb Z18788	
1302	AF100694	complete cds	le-28	4056454	come from this gene.	
			1 10 20	4030434	[Arabidopsis thaliana]	1e-09
	İ	1	1 1		(AC005990) Contains repeated region with similarity to	
	f		1 1		gb U43627 extensin (atExt1)	
		-	1 1		gene from Arabidopsis thaliana.	
		Mus musculus	1 1		ESTs gb Z34165 and gb Z18788	
		Pontin52 mRNA,	1 1	4	come from this gene.	
303	AF100694	complete cds	le-28	4056454	[Arabidopsis thaliana]	
ı					(AC005990) Contains repeated	9e-10
- 1			1 1		region with similarity to	
- 1	-		1 1		gb U43627 exiensin (atExt1)	
1			1		gene from Arabidopsis thaliana.	
		Mus musculus			ESTs gb Z34165 and gb Z18788	
304		Pontin52 mRNA,	1 1		come from this gene.	
	AF100694	complete cds	1e-28	4056454	[Arabidopsis thaliana]	4e-10
		·	, T		(AC005990) Contains repeated	46-10
ı] [•	region with similarity to	
ł	ł		1 1		gb U43627 extensin (atExt1)	
-	l,	Aus musculus		1	gene from Arabidopsis thaliana.	
J		Pontin52 mRNA,	1		ESTs gb Z34165 and gb Z18788	
05		omplete cds	, ,		come from this gene.	
1		ompiete cus	1e-28	4056454	[Arabidopsis thaliana]	9e-11
ı	1		1		(AC005990) Contains repeated	
	1	ľ	1		region with similarity to	
- 1	- 1.		1		gb U43627 extensin (atExt1)	j
	l _N	lus musculus	J].	gene from Arabidopsis thaliana.	
ı		ontin52 mRNA,	j	Į:	ESTs gb Z34165 and gb Z18788	
06		omplete cds	10.20	10551-	come from this gene.	İ
			1e-28	4056454	[Arabidopsis thaliana]	6e-11

<u> </u>		st Neighbor (BlastN vs. (Genbank)	Nearest Neis	hbor (BlastX vs. Non-Redundant P	-atains)
SE	2			1	13. Non-Redundant P	roteins)
ID	ACCESSIO	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	DVALLE
_		Mus musculus			J J J J J J J J J J J J J J J J J J J	P VALU
1.20		Pontin52 mRNA.				
130	7 AF100694	complete cds	4e-29	<none></none>	<none></none>	<none:< td=""></none:<>
130	8 AF079529	Homo sapiens cAMP specific phosphodies terase 8B	I	No.		
	120,732	phosphodiesterase an	4e-29	<none></none>	<none></none>	<none></none>
		H.sapiens mitochondrial DNA,			CYTOCHROME COXIDASE POLYPEPTIDE I chain I - human mitochondrion (SGC1) >gi 13006 (V00662) cytochrome oxidase I [Homo sapiens] >gi 506829 (J01415) cytochrome oxidase subunit 1	
1309	X93334	complete genome	4e-29	1 16977	[Homo sapiens] sapiens]	3e-09
1310	AF020760	Homo sapiens serine protease (Omi) mRNA, complete cds	4e-29	2530017	(AF020760) serine protease	30-09
		Xenopus laevis	46-29	2738915	[Homo sapiens]	8e-12
1311	U95097	mitotic phosphoprotein 43 mRNA, partial cds Homo sapiens	4e-29	2072294	(U95097) mitotic phosphoprotein 43 [Xenopus laevis]	le-25
1312	L32162	transcription factor mRNA, 5' end.	2e-29	2501706	RENAL TRANSCRIPTION FACTOR KID-1 finger protein [Mus musculus]	8e-15
1313	AE100604	Mus musculus Pontin52 mRNA,			(AC005990) Contains repeated region with similarity to gb U43627 extensin (atExt1) gene from Arabidopsis thaliana. ESTs gb Z34165 and gb Z18788 come from this gene.	
.515	AF100694	complete cds	1e-29	4056454	[Arabidopsis thaliana]	1e-04
314	AF100694	Mus musculus Pontin52 mRNA, complete cds	Ie-29		FMRFAMIDE-RELATED NEUROPEPTIDES PRECURSOR >gi 416208 (U03137) neuropeptide precursor FMRFamide-related	
		Homo sapiens g16 protein (g16) mRNA,	10-29		peptide [Lymnaea stagnalis] (AF069517) RNA binding	1e-05
315		complete cds	le-29		protein DEF-3 [Homo sapiens]	6e-10

	Nearest l	Veighbor (BlastN vs. G	enbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant Pr	oteins)
SEQ						
ΙD	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
					intercellular adhesion molecule	
					3. ICAM-3=lymphocyte	
					function-associated antigen 1	
i I					counter-receptor homolog	
1216	V40711	H. sapiens mRNA for	5- 20	200256	[human, tonsil, Peptide Partial,	2.00
1316	X69711	ICAM-R Homo sapiens	5e-30	299356	518 aa]	3e-08
		receptor-associated			(AF012108) Amplified in Breast	İ
1317	AF010227	coactivator 3	5e-30	2331250	Cancer [Homo sapiens]	8e-09
1317	AI 010227	Homo sapiens full	36-30	2331230	Cancer (Hollio sapiens)	00-03
		length insert cDNA			(AJ235273) CELL SURFACE	
1318	AF086395	clone ZD75C01	2e-30	3861241	ANTIGEN (sca5)	4.2
			· .		PHOSPHOGLYCERATE	
					KINASE 2.7.2.3) - Pyrococcus	
		Human 28S			woesei >gi 1054832 (X73527)	
		ribosomal RNA gene,			phosphoglycerate kinase	
1319	M27830	complete cds.	2e-30	1730522	[Pyrococcus woesei]	3.8
		M. Comparison			RAS-RELATED PROTEIN	
		Mouse GTP-binding			RAB-17 Rab17 - mouse	
1320	M79307	protein (Rab17) mRNA sequence.	2e-30	464564	(fragment) >gi 297157 (X70804) rab17 [Mus musculus]	9e-11
1320	19307	Human DNA	2e-30	404304	(X70804) Fab 17 [Mus musculus]	96-11
		sequence from clone				
		U247E12 on	ļ			
		chromosome Xq22-	i			
		23, complete				
		sequence [Homo			(U93570) putative p150 [Homo	
1321	AL022168	sapiens]	1e-30	2072967	sapiens]	3e-11
		M.musculus pacsin	_ (<u>.</u> j			
1322	X85124	gene	le-30	2217964	(Z50798) p52 [Gallus gallus] structural polyprotein -	le-34
			1		Venezuelan equine encephalitis	
					virus (strain TRD) >gi[323710	
	ļ	Homo sapiens	ļ	• •	(J04332) poly-envelope protein	
		phosphoprotein CtBP			[Venezuelan equine encephalitis	
1323	U37408	mRNA, complete cds	5e-31	74518	virus1	1.1
		Human lens		, ,,,,,,		
ŀ		membrane protein	j			
		(mp19) gene, exon	l		!!!! ALU SUBFAMILY J	
1324	L04193	11.	2e-31	728831	WARNING ENTRY	7e-07
				•		
		Human 28S	1		1	
1325	M11167	ribosomal RNA gene.	6e-32	<none></none>	<none></none>	<none></none>

	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neigh	bor (BlastX vs. Non-Redundant Pi	roteins)
SEQ						
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
						1
		Human cAMP-				†
1		dependent protein		<u>[</u>	1]
1		kinase type I-alpha		Ī		1
1	İ	subunit (PRKARIA)				
1326	M33336	mRNA, complete cds	2e-32	<none></none>	<none></none>	<none></none>
		Human				
		glucocerebrosidase]	1	1
1,,,,,	702040	pseudogene, complete			glucosylceramidase (EC	1
1327	J03060	cds Human lipid-	2e-32	2144479	3.2.1.45) precursor - human	le-05
					·	
ł	\	activated protein kinase PRK1 mRNA,				
1328	U33053	complete cds	7e-33	2137690	protein kinase (EC 2.7.1.37) -	
1328	60000	complete cas	/6-27	2137689	mouse	le-14
		Human elongation				1
]		factor EF-1-alpha				
1 1		gene, complete cds. >				1
ļ ļ		:: dbj E02629 E02629				
		DNA of human				
		polypeptide chain				1
		elongation factor- l				
1329	J04617	alpha	6e-33	<none></none>	<none></none>	<none></none>
		Homo sapiens (clone s22i71) mRNA			INTERMEDIATE FILAMENT	
1330	L40396	fragment	6e-33	124225	PROTEIN B protein B -	
1330		S.cerevisiae	06-33	124235	common_roundworm	1.00
i I		chromosome VII	i		MSPI PROTEIN HOMOLOG]
	,	reading frame ORF	Í		Yeast MSP1 protein (TAT-	1
1331		YGR028w	6e-33	1709135	binding homolog 4)	8e-50
		Homo sapiens mRNA	İ		(U42471) Wiscott-Aldrich	
	1	for KIAA0472 .	- 1		Syndrome protein homolog	
1332	AB007941	protein, partial cds	2e-33	1150834	[Mus musculus]	2.0
	1.	Pattus nonuscious				
		Rattus norvegicus putative peroxisomal				
ŀ		2,4-dienoyl-CoA]	
		reductase (DCR-	Í		(A E044574) purativa	
		AKL) mRNA,			(AF044574) putative	
1333		complete cds	2e-34	4105269	peroxisomal 2,4-dienoyl-CoA reductase [Rattus norvegicus]	6e-15
		Human mRNA for		4103209	readerase [rearras noivegicus]	06-13
		KIAA0101 gene,				
1334		complete cds	7 e-3 5	<none></none>	<none></none>	<none></none>
					trithorax homolog HTX - human	
]		H.sapiens p63 mRNA	l l		(fragment) homolog=MILL	
	•	for transmembrane		•	{alternative splicing, clone 14p-	1
1335	X69910 F	protein	7e-35	2136323	18B}	0.94

art.	Nearest l	Neighbor (BlastN vs. G	enbank)	Nearest Neight	oor (BlastX vs. Non-Redundant Pr	oteins)
SEQ						
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Homo sapiens				
		tetraspan TM4SF				
		(TSPAN-5) gene,			(AF065389) tetraspan NET-4	
1336	AF053455	complete cds	7e-35	3152703	[Homo sapiens]	le-25
1337	V50774	D.melanogaster crn mRNA	3e-35	117478	CROOKED NECK PROTEIN	6e-41
133/	X58374	Homo sapiens full	36-35	11/4/8	CROOKED NECK PROTEIN	06-41
		length insert cDNA			(AF031328) aminoglycoside 6'-	
1338	AF086492	clone ZD95D11	9e-36	2909809	N-acetyltransferase It	1.9
		H.sapiens telomeric				
		DNA sequence, clone 12PTEL 120, read		<i>'</i> '		
1339	Z96223	12PTELOO120.seq	3 e-36	2408068	(Z99165) hypothetical protein	0.61
		H.sapiens mRNA for				
		phenylalkylamine			emopamil-binding protein -	
1340	Z37986	binding protein.	1e-36	· 1362793	human >gi 780263	5e-11
		Human ribosomal				
		protein S27 mRNA.		•		
		complete cds. end				
! !		similar to similar to				
1 1		metallopanstimulin 1		<u>.</u>		
		> :: - LIA A 2 I 6 2 2 7 I A A 2 I 6	•			1
		gb AA316327 AA316 327 EST188061 HCC			40S RIBOSOMAL PROTEIN	
		cell line (matastasis to			S27 growth factor-inducible zinc	
		liver in mouse) [[finger protein MPS-1 - human	
		Homo sapiens cDNA			>gi 431319 (L19739)	
		5' end similar to			metallopanstimulin [Homo	
		similar to			sapiens] >gi 1373421 (U57847)	
1341	U57847	metallopanstimulin l	3e-37	1171014	ribosomal protein S27	1.4
		Dattus paragiaus	,		70 KD WD-REPEAT TUMOR- SPECIFIC ANTIGEN	
		Rattus norvegicus mRNA for 70 kDa	ı		>gi 2505957 gn1 PID e353992	
		tumor specific			(Y15054) 70 kD tumor-specific	
1342		antigen, partial	3e-37	3123027	antigen [Rattus norvegicus]	2e-15
		Rattus norvegicus serine/threonine			(AF084205) serine/threonine	
		protein kinase TAO1	ł		protein kinuse TAOI [Rattus	
1343		mRNA, complete cds	3e-37	3452473	norvegicus]	5e-4"
.5 75		R.norvegicus	30 31	J 132173	No Optodo	
		(Sprague Dawley)	İ			
		ARL5 mRNA for				
1344	X78604	ARF-like protein 5	le-37	<none></none>	<none></none>	<none></none>

- 7	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ						
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Homo sapiens				
		chromosome 22 CpG			,	
		island DNA, genomic				
		Mse1 fragment, clone	·		1	
		22CGIB49A3 .			I	
1345	AJ236644	complete read	1e-37	2239219	(Z97210) hypothetical protein	6e-05
ا ا	*****	Human zinc finger	4 20	2127270	DNA-binding protein - mouse	7. 22
1346	U09367	protein ZNF136	4e-39	2137269	>gi 437444	7e-23
		Human DNA				
1 1		sequence from		,		
1 1		cosmid L69F7B,				
		Huntington's Disease	• •		1	
		Region, chromosome				!!!
		4p16.3 contains			(AL023094) putative cyclase	1
		Huntington Disease			associated protein CAP	1
1347	Z69649	(HD) gene.	3e-39	3096918	[Arabidopsis thaliana]	5.6
-						
		Homo sapiens				
li		tetraspan NET-4		·	(AF065389) tetraspan NET-4	
1348	AF065389	mRNA, complete cds	1e-39	3152703	[Homo sapiens]	6e-29
		Homo sapiens clone				
		23923 mRNA			(U60883) CapC (Bacillus	
1349	AF038172	sequence	1e-40	1813464	firmus]	2.8
		H sapiens Fanconi				
		anaemia group A			-i firman	}
1,250	702005	gene, exons 39, 40,	1. 40	2127070	zinc finger protein - mouse	3e-23
1350	Z83095	41, 42 and 43 Homo sapiens 17-	le-40	2137870	(fragment)	36-23
		beta-hydroxysteroid				i l
	i	dehydrogenase IV			(AL008730) dJ487J7.1.1	} i
		(HSD17B4) gene,			(putative protein dJ487J7.1	
1351	AF057734	exon 16	1e-40	2842416	isoform 1) [Homo sapiens]	6e-61
	214 05/154	Homo sapiens clone	10.10	20.2710		
		24544 beta-				
		dystrobrevin mRNA.			(Y15718) dystrobrevin B DTN-	
1352	AF070567	partial cds	4e-41	3133087	B2 [Homo sapiens]	7e-13
	·	Homo sapiens Arp2/3				
		protein complex				
		subunit p16-Arc	l			
	_	(ARC16) mRNA,			ARP2/3 COMPLEX 16 KD	
1353	AF006088	complete cds	2c-41	3121767	SUBUNIT	3e-36
]		M.musculus mRNA			(45016110) No. 4-5-14	
,,,,,	W/0010	of enhancer-trap-	6- 43	2201152	(AF016418) No definition line] , [
1354	X69942	locus l	6e-42	2291152	found [Caenorhabditis elegans]	6.4

· ·	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
מו	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		H.sapiens mRNA for			(U28811) cysteine-rich		
1355	X87838	beta-catenin	5e-42	1373019	fibroblast growth factor receptor	8c-05	
						i	
		Homo sapiens mRNA for KIAA0725			(AB018268) KIAA0725 protein		
1356	AB018268	protein, partial cds	5e-42	3882171	[Homo sapiens]	2e-33	
1330	AD010200	Human cathepsin E	30-42	3002171	[Liono suprens]	20 33	
		(CTSE) gene, exon 9					
1357	M84424	and complete cds.	2e-42	<none></none>	<none></none>	<none></none>	
					(U80776) ort; encodes putative		
		TY FOT -I			chimeric protein with SET domain in N-terminus with		
		Human EST clone NIB1543 mariner			similarity to several other		
		transposon Hsmarl			human, Drosophila, nematode		
		orf gene, complete			and yeast proteins [Homo		
1358	U80776	cds	2e-42	223 1380	sapiens]	3e-11	
		Human G protein					
		Golf alpha gene, exon			(AF067608) No definition line		
1359	U55184	12 and complete cds Homo sapiens PAC	2e-42	3165531	found [Caenorhabditis elegans]	1e-16	
		clone DJ1152D16					
		from Xq23, complete					
	•	sequence [Homo			(AB007407) myeloid zinc finger		
1360	AC005190	sapiens]	6e-43	2978255	protein-2 [Mus musculus]	2.3	
] [Homo sapiens mRNA					
1261	A DO10204	for KIAA0741	52.42	NONE	<none></none>	<none></none>	
1361	AB018284	protein, complete cds	5e-43	<none></none>	SAGIVES	CHOILE	
		Homo sapiens mRNA	ĺ				
		for KIAA0565	,		(AB011137) KIAA0565 protein		
1362		protein, complete cds	5e-43	3043654	[Homo sapiens]	1e-07	
		Human set gene,					
1363	M93651	complete cds.	2e-43	<none></none>	<none></none>	<none></none>	
					(U84404) E6-associated protein		
	ļ	H.sapiens mRNA for	İ		E6-AP/ubiquitin-protein ligase		
		RNA polymerase II			[Homo sapiens] >gi 2361031		
		elongation factor-like			(AF016708) E6-AP ubiquitin-	·	
1364	Z47087	protein.	2e-43	1872514	protein ligase [Homo sapiens]	7.2	
		Drosophila					
		melanogaster pelota	ł		PELOTA PROTEIN >gi 973224		
1265	1	(pelo) mRNA.	20.42	1252726	(U27197) pelota [Drosophila melanogaster]	le-46	
1365	U27197	complete cds	2e-43	1352736	Imeranogasteri	16-40	

25, 25	Nearest Neighbor (BlastN vs. Genbank)		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ						
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
					RRP5 PROTEIN HOMOLOG	
		Human mRNA for			(KIAA0185) hypothetical	
		KIAA0185 gene.			protein YM9959.11C of	
1366	D80007	partial cds	·6e-44	2498864	S.cerevisiae. [Homo sapiens]	6e-09
		Homo sapiens				l i
		secretory carrier			1	<u> </u>
		membrane protein			(AF005039) secretory carrier]
1000	. ======	(SCAMP3) mRNA,		0000010	membrane protein [Homo	2-00
1367	AF005039	complete cds	6e-44	2232243	sapiens] (X68101) trg gene product	2e-09
1368	X68101	R.norvegicus trg mRNA	2e-44	550420	[Rattus norvegicus]	1e-37
1300	V00101	Homo sapiens	26-44	330420	[Kattas not vegicus]	10-57
		cyclooxygenase				
1		(COX-2) gene,	••		(U93565) putative p150 [Homo	
1369	AF044206	promoter and exon 1	2e-45	2072953	sapiens)	5e-06
		Homo sapiens				
		faciogenital dysplasia				
		(FGD1) gene, 5' end				
1370	L48708	of intron 17	8e-46	<none></none>	<none></none>	<none></none>
					POLYPEPTIDE VIIA-LIVER	
					PRECURSOR	}
					>gi 2144370 pir OSHU7L	
i I					cytochrome-c oxidase (EC	
		Human COX VIIa-L			1.9.3.1) chain VIIa precursor.	
		mRNA for liver-			hepatic - human >gi 30147	
		specific cytochrome c			(X15822) precursor (AA -23 to	
1371		oxidase (EC 1.9.3.1.)	3e-46	117121	60) [Homo sapiens]	5e-13
		, , , , , , , , , , , , , , , , , , , ,				
		Mus musculus		ı		
] !		stromal cell protein			(U47323) stromal cell protein	1
1372	U47323	mRNA, complete cds	3c-46	1493833	[Mus musculus]	le-48
ļ . T				•	PROTEIN T28D9.7 IN	
	•	·			CHROMOSOME II >gi 861264	
i i	į				(U28738) coded for by C.	
					elegans cDNA yk8h5.3; coded	
			i		for by C. elegans cDNA	
		Homo sapiens			yk8h5.5; similar to C. elegans	
		reticulon gene family			deg-1 and mec-4 in exon 2	
1373		protein	7e-47	1731169	[Caenorhabditis elegans]	7.8
13,2	.11 033327	Homo sapiens mRNA			Table 1	
{		for puromycin	1			
		sensitive				
		aminopeptidase,			(U39123) T cell receptor beta	
1374		partial	3e-47	1777519	chain [Homo sapiens]	9.7

	Nearest l	Nearest Neighbor (BlastN vs. Genbank)		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
						 	
		Homo sapiens histone	-		(U83821) NADH		
		binding protein			dehydrogenase subunit 3		
1375	M97856	mRNA, complete cds.	3e-47	2645327	[Oryzomys palustris]	5.7	
		Human retinoblastoma-			C) D CIALIGACID		
		related Rb2/p130			CMP-SIALIC ACID		
		gene, 5' flanking			TRANSPORTER CMP-sialic		
1376	U53220	region and partial eds	3c-47	2499225	acid transporter [Cricetulus griseus]	5.3	
1370	033220	H.sapiens mRNA for	36-47	2499223	griscus	3.3	
		hepatocyte nuclear			!!!! ALU SUBFAMILY SB	1	
1377	X87870	factor 4a	1e-47	728832	WARNING ENTRY	7.3	
	33970					 	
		Mus musculus		•			
	·	proteasome regulator				l · .	
		PA28 beta subunit					
1378	AF060195	gene, complete cds	3e-48	478681	limb deformity protein - chicken	0.25	
					TESTIS SPECIFIC PROTEIN		
ı					A (ZINC FINGER PROTEIN		
					TSGA) >gi 281040 pir S28499	1	
		Homo sapiens mRNA			probable zinc finger protein - rat		
1270	47010006	for KIAA0742		2122262	>gi 57504 (X59993) zinc finger		
1379		protein, partial cds Human endogenous	1e-48	3122969	protein	1e-30	
ŀ		retrovirus clone					
I		c5.11, HERV-H					
f	ļ	multiply spliced					
		subgenomic leader,					
		protease and integrase	ŀ				
		region mRNA, partial			retroviral proteinase-like protein		
1380		cds	4e-49	88558	- human	6e-05	
ľ		Homo sapiens			·		
ļ		mRNA, chromosome	Į				
	1	l specific transcript	. i				
1381	AB007956	KIAA0487	1e-49	<none></none>	<none></none>	<none></none>	
ı	1		1		ALPHA SCRUIN >gi 633238 (Z38132) scruin [Limulus		
- 1	1	Homo sapiens mRNA			T -		
ł	1	for KIAA0214	·		polyphemus] >gi 1093326 prf 2103269A		
1382		protein, complete cds	le-49	2497944	Scrulin (Limulus sp.)	9.7	
-552		Human transcription	10-49	Z47/744	Securit (Limitus Sp.)		
		factor (SC1) gene,	Į				
1383		complete cds.	4e-50	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEO			1			1	
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		Mus musculus ATP-				Î ·	
		dependent RNA					
1		helicase mRNA,			(U46690) ATP-dependent RNA	1 .	
1384	U46690	partial cds.	4e-50	1335873	helicase [Mus musculus]	3e-24	
		Mus musculus					
		claudin-2 mRNA,			(AF072128) claudin-2 [Mus		
1385	AF072128	complete cds	2e-50	3335184	musculus]	4e-24	
		Homo sapiens					
		snRNA activating		1	•		
		protein complex		•			
		19kDa subunit			(AF093593) snRNA activating		
		(SNAP19) mRNA,	1	• •	protein complex 19kDa subunit		
1386	AF093593	complete cds	le-50	3668416	[Homo sapiens]	0.003	
		Homo sapiens			·		
		monocarboxylate					
		transporter					
		homologue MCT6			(X92485) pva I [Plasmodium		
1387	U79745	mRNA, complete cds	le-50	1177607	vivax]	2e-07	
		Rattus norvegicus					
		hepatocyte nuclear			(L10409) fork head related		
1388	L09647	factor 3a	1e-50	404764	protein [Mus musculus]	2e-21	
		Mouse E46 mRNA					
1389	X61506	for E46 protein	4e-51	114909	BRAIN PROTEIN E46	1e-20	
1 1					LINE-I REVERSE		
1 1		Human debrisoquine			TRANSCRIPTASE	1	
i 1		4-hydroxylase			HOMOLOG protein	i	
1390	M33387	(CYP2D8P) and	1e-51	126296	[Nycticebus coucang]	5e-15	
ll			1				
ĺĺ		Homo sapiens zinc					
i I		finger protein (ZPR1)	1			•	
1391	AF019767	mRNA, complete cds	4e-52	961507	(D63788) anchor protein, LCM	5.9	
1			ľ	• ()			
1 1		H.sapiens mRNA for	J			i	
. <u>.</u>		phenylalkylamine	. 1				
1392	Z37986	binding protein.	2e-52	<none></none>	<none></none>	<none></none>	
1	l		ļ		(Z49128) weak similarity with	1	
1	Į	ĺ	· 1		SINR protein (Swiss Prot		
	[į		accession number P06533);	•	
	j		ļ		cDNA EST EMBL:T00631		
[l.	VI	j		comes from this gene; cDNA		
	- 1	Human MHC class 1			EST yk293d10.5 comes from		
1303		molecule (MICB)			this gene [Caenorhabditis	ľ	
1393	U65416	gene, complete cds	2e-52	3878637	elegans)	8.7	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	1.52.034			TACTICS! IACIBL	Diasta vs. Non-Regundant Pi	(Oterns)	
ID	ACCESSION	DESCRIPTION	BVALUE	A CCERCION	DESCRIPTION.		
-	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					beta-globin DNA-binding		
					protein B1, transcription factor	1	
					PU.1 - mouse >gi 200586	1	
		H.sapiens CpG DNA,			(M32370) PU.1 protein [Mus		
		clone 189a6, forward			musculus) >gi 200972		
1394	Z57647	read cpg189a6.ft1a.	2e-52	111187	(M38252) transcription factor Pu.1 [Mus musculus]		
1321	257047	Human activated	26-32	11310/	Fu.1 [Mus musculus]	5.8	
1 1		p21cdc42Hs kinase			(AF037260) non-receptor	}	
1 1		(ack) mRNA.			protein tyrosine kinase Ack	i	
1395	L13738	complete cds.	2e-52	2921447	[Mus musculus]	7e-23	
		o inprote des	20 32	. 2721477	[17103 Museurus]	76-23	
						İ	
		Homo sapiens spindle				Į	
		pole body protein	ŀ			<u> </u>	
i 1		spc97 homolog GCP2	1		(AF042379) spindle pole body	1	
1396	AF042379	mRNA, complete cds	7e-53	2801701	protein spc97 homolog GCP2	le-16	
					DNA-DIRECTED RNA		
}			ł		POLYMERASE I 40 KD		
					POLYPEPTIDE (RPA40)		
		Homo sapiens RNA	1		(RPA39) >gi 2266929]	
		polymerase I 40kD			(AF008442) RNA polymerase I		
1207	. =0 . =	subunit mRNA,			subunit hRPA39 [Homo		
1397		complete cds Homo sapiens cell	6e-53	3914807	sapiens]	4e-19	
		-	1			j	
		cycle protein (PA2G4) gene, exons	Į.			i	
l		6 through 13, and	1				
1398		complete cds	2e-53	<none></none>	NONE	ALONIES	
1370	14 10-070	complete cus	26-33	CHONES	<none></none>	<none></none>	
]	{ VNTR locus DXZ4,					
1	Ĭ	hypervariable tandem	i			1	
		repeat cluster}					
I	Ì	[human, Genomic,	·	•			
- 1	ł	2991 nt] > ::			(U38810) CAGR1 [Homo		
- 1].	gb L07935 HUMVNT	j		[sapiens] >gi 3098420		
- 1		RA Homo sapiens	ļ		(AF040945) homeotic regulator		
1	ļ	microsatellite VNTR	[homolog MAB21 [Mus		
1399		DNA sequence.	2e-53	1209669	musculus]	4.6	
		Human mRNA for					
		KIAA0218 gene,	ľ		(AC005168) unknown protein		
1400	D86972	complete cds	1e-53	3426041	[Arabidopsis thaliana]	9.1	

PCT/US00/18374

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (Blast V. vo. New D. J. J. D. V.			
SEQ		1	T T	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
1D	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUI	
<u> </u>							
1401	AJ236682	Homo sapiens chromosome 22 CpG island DNA, genomic Mse1 fragment, clone 22CG1B49E6, complete read	7e-54	3928721	(AL034355) putative cytochrome oxidase subunit I [Streptomyces coelicolor]	0.30	
1402	AJ236682	Homo sapiens chromosome 22 CpG island DNA, genomic Msel fragment, clone 22CGIB49E6, complete read	6e-54	3928721	(AL034355) putative cytochrome oxidase subunit I [Streptomyces coelicolor]	0.28	
		Human histone					
1403	M37583	(H2A.Z) mRNA, complete cds.	6e-54	70711	histone H2A.F, embryonic - chicken	2e-16	
1404	AJ009947	Homo sapiens mRNA for putative ATPase. partial	6e-54	3550295	(AJ009947) putative ATPase [Homo sapiens]	3e-18	
		B.taurus mRNA for			[2235 0-21.0.13]	, 30-10	
1405	Y08459	novel cytoplasmic protein	2e-54	<none></none>	<none></none>	<none></none>	
1406		Homo sapiens BC-2 protein mRNA, complete cds	2e-54	2828147	(AF042384) BC-2 protein [Homo sapiens]	2e-14	
1407	AF042379	Homo sapiens spindle pole body protein spc97 homolog GCP2 mRNA, complete cds	8e-55	2801701	(AF042379) spindle pole body protein spc97 homolog GCP2	2e-17	
1408	i	Oryctolagus cuniculus translation initiation factor eIF2C mRNA, complete cds	7e-55	3253159	(AF005355) translation	2. 52	
]]	Homo sapiens RNA polymerase I subunit IRPA39 mRNA,	7,0-33	3233139	initiation factor elF2C (AF047441) RNA polymerase 1	3e-53	
1409		omplete cds	3e-55	3335138	40kD subunit [Homo sapiens]	3c-20	
1410	F s	Homo sapiens RNA polymerase I 40kD ubunit mRNA, omplete cds	3e-55		(AF047441) RNA polymerase I		
	07/471 10	ompiete cus	36-33	3335138	40kD subunit [Homo sapiens]	3e- <u>20</u>	

<u> </u>	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ	1							
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
		Human mRNA for						
		Rap1B protein > ::						
1		emb A08693 A08693		•				
		H.sapiens rap 1 b			transforming protein rap1b - rat			
1411	X08004	cDNA	2e-55	539995	(strain Copenhagen)	2e-18		
1	1							
		Homo sapiens ALR			(AF010403) ALR [Homo			
1412	AF010403	mRNA, complete cds	2e-55	2358285	sapiens]	1e-49		
	1	Human transmodulin			(SE2010) and 5' of Each (miss			
1413	M77016	Human tropomodulin mRNA, complete cds.	8e-56	262249	(S52010) orf1 5' of EpoR [mice, Peptide, 85 aa] [Mus sp.]	0.027		
1713	14177010	miktyA, complete cus.	86-30	202249	Peptide, 83 aaj [ivius sp.]	0.027		
		Homo sapiens mRNA	٠,		1			
ł		for KIAA0826			1			
1414	AB020633	protein, partial cds	2e-56	<none></none>	<none></none>	<none></none>		
		H.sapiens genomic			7	4,01,122		
		DNA (chromosome			(U84501) cuticle collagen			
1415	X87489	3: clone NL1243D)	2e-56 [.]	1814029	[Caenorhabditis briggsae]	0.038		
		Homo sapiens						
		KIAA0433 mRNA,			(AB007893) KIAA0433 [Homo			
1416	AB007893	partial cds	2e-56	2887437	sapiens]	9e-21		
! i		H.sapiens HZF2	1	•	(AF054180) hematopoietic cell			
ا ـ ا		mRNA for zinc finger			derived zinc finger protein			
1417	X78925	protein	1e-56	3342002	[Homo sapiens]	2e-21		
i I		H.sapiens mRNA for interferon regulatory			INTERFERON			
1418	Z56281	factor 3	9e-57	2407442	REGULATORY FACTOR 3	2e-21		
1410	230281	Homo sapiens nuclear	96-37	2497442	factor 3 [Homo sapiens]	26-21		
		VCP-like protein	i		(U68140) nuclear VCP-like			
1419	U78772	NVLp.1	8e-57	2406565	protein NVLp.2 [Homo sapiens]	5e-20		
		Human mRNA for			(D79994) similar to ankyrin of			
		KIAA0172 gene,			Chromatium vinosum. [Homo			
1420	D79994	partial cds	3e-57	1136404	sapiens]	9e-38		
	1	Human mRNA for						
	1	KIAA0344 gene,	1		(AB002342) KIAA0344 [Homo			
1421	AB002342	complete cds	1e-57	2224629	sapiens]	4c-20		
			T					
			1					
l		Human transaldolase	1					
I		mRNA containing	- 1		(1/2150) · · · · · · · · · · · · · · · · · · ·			
1422		transposable element,	, 57	1553110	(U63159) transaldolase [Mus			
1422	L19437	complete cds	1e-57	1553119	musculus] PROBABLE ATP-	2e-20		
			1		DEPENDENT RNA			
· [i		HELICASE P54 (ONCOGENE			
Į	1	Human mRNA for		•	RCK) (DEAD BOX PROTEIN			
1423		RCK. complete cds	9e-58	129376	(6)	le-10		
1					I 🗸 ,	10 10		

£72414	Nearest	Neighbor (BlastN vs. C	Genbank)	Nearest Neigl	hbor (BlastX vs. Non-Redundant Pr	oteins)
SEC			1	1.02.03.1.01g.	Ton-Redundant 1	T T
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
-	 	11				
	j	H.sapiens BDPI	İ		1	
1424	X79568	mRNA for protein-	0.50		(X79568) protein-tyrosine-	
1727	X/9308	tyrosine-phosphatase	9e-58	1871531	phosphatase	1e-22
į		H.sapiens BDP1				
ľ		mRNA for protein-	i i		(X79568) protein-tyrosine-	1
1425	X79568	tyrosine-phosphatase	9e-58	1871531	phosphatase	9e-23
		Homo sapiens	3000	1071331	phosphatasc	96-23
		HKE1.5 mRNA for				
1	j .	GDS-related protein,				1 1
1426	AB012295	complete cds	7e-58	2648021	(Z97184) RGL2 [Homo sapiens]	9e-19
1	1		• -		glutamine (Q)-rich factor 1,	
1		Homo sapiens full			QRF-1 - mouse factor 1, QRF-1	
1,407	150000	length insert cDNA			[mice, B-cell leukemia, BCL1,	
1427	AF086040	clone YX52E07 Homo sapiens ca xi	. 1e-58	543222	Peptide Partial, 84 aa]	3e-36
1		mRNA for carbonic				
1		anhydrase-related				1
1		protein XI, complete			·	
1428	AB018195	cds	4e-59	<none></none>	<none></none>	NONE
		Mus musculus IRE1		- CHOILD	NONE>	<none></none>
		(Ire1) mRNA,			(AF071777) IREI [Mus	
1429	AF071777	complete cds	4e-59	3766209	musculus]	7e-28
l]		Homo sapiens mRNA				l
}		for SH3 binding				i
1430	4 D000440	protein, complete cds,	[1	ı
1430	AB000462	clone:RES4-23A Homo sapiens clone	3e-59	<none></none>	<none></none>	<none></none>
		23923 mRNA			(TORES IN NAVIONALIA	1
1431	AF038172	sequence	3e-59	3758855	(Z98551) MAL3P6.11	
	112 55 5.7.2	Human DNA			[Plasmodium falciparum]	1.3
	•	sequence from phage	ľ		·	
		pTEL from a contig			1	1
		from the tip of the		•	1	
	l l	short arm of]	1
1	T I	chromosome 16,			ļ	1
1		spanning 2Mb of	1		RIBONUCLEOPROTEIN	
1430		16p13.3 Contains	Ĭ	,-	RB97D ribonucleoprotein	
1432	Z84812	ESTs	1e-59	400927	[Drosophila melanogaster]	2.5
- [Human laminin-	ł			
1		binding protein gene.				
ļ		partial cds, and E2	l		İ	į
I	L T	small nucleolar RNA				
- 1		gene, complete		i	·.	
1433	4	sequence	1e-59	226005	protein 40kD [Mus musculus]	7e-05
					(c. cas mascaras)	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ						1	
ΙD	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					DUAL SPECIFICITY		
1					MITOGEN-ACTIVATED	,	
1					PROTEIN KINASE KINASE 2		
		Homosapiens ERK	ĺ	İ	(MAP KINASE KINASE 2)	1	
1	1	activator kinase		Į	(MAPKK 2) kinase type 2	· ·	
1434	L11285	(MEK2) mRNA.	le-59	2499630	[Gallus gallus]	3e-21	
ł	i	Homo sapiens full					
1	l	length insert cDNA	ĺ	·	(AC005239) F23149_1 [Homo		
1435	AF086555	clone ZE14E04	4e-60	3287674	sapiens]	2e-04	
1	Ī	Human (clone					
i		pHAIV2-12) alpha-2			(X05610) alpha (2) chain	1.	
1436	M24766	collagen type IV	4e-60	29551	[Homo sapiens]	6e-15	
1	i	H.sapiens mki67a	• .		ANTIGEN KI-67		
ı	ł	mRNA (long type)			>gi 539555 pir A48666 cell		
		for antigen of			proliferation antigen Ki-67, long	1	
		monoclonal antibody	·		form - human Ki-67 [Homo	Ì	
1437	X65550	Ki-67	4e-60	1170654	sapiens]	3e-15	
l					(X05949) Calmodulin (AA 2 -		
		Human calmodulin			59) (449 is 1st base in codon)		
1438	M27319	mRNA, complete cds.	4e-60	1345451	[Drosophila melanogaster]	7e-20	
i .							
		Homo sapiens mRNA	1				
l i		for transducin (beta)			(X06172) put. 134 kD protein	i	
1439	Y12781	like I protein	3e-60	62133	(AA 1 - I 187); put. replicase	7.4	
		Human mRNA for					
		KIAA0385 gene.	[•		
1440	AB002383	complete cds	1e-60	1001548	(D64000) hypothetical protein	4.4	
		Homo sapiens clone	ł				
		24732 unknown			(AF070614) unknown [Homo		
1441	AF070614	mRNA, partial cds	2e-61	3283879	sapiens]	3e-17	
ı			- 1	•	MICROTUBULE-		
		Human mRNA for			ASSOCIATED PROTEIN 4		
		KIAA0328 gene.		•	microtubule-associated protein-		
1442		partial cds	6e-62	547891	U [Bos taurus]	5.6	
		Homo sapiens full	1		1		
1443		length insert cDNA]		
1443	AF086471	clone ZD88A01	5e-62	<none></none>	<none></none>	<none></none>	
1			Ì		DIHYDROXYPHENYLPROPI	-	
ļ	Ī		Į.		ONATE 1.2-DIOXYGENASE		
- 1	ļ	İ	.]			·	
i	1		i		>gi 1657544 (U73857) similar		
- 1	ĺ		}		to mcpl gene (catechol 2.3-		
J	ĺ]	1		dioxygenuse) of A. eutrophus 3-		
	į		1		(2,3-	į	
- 1	l,	Human mRNA for	i		dihydroxyphenylpropionate)1, 2-	ļ	
1	1		J		dioxygenuse 2.3-	į	
1444		KIAA0313 gene,	2 (2	0506555	dihydroxyphenylpropionate 1,2-		
7 774	AB002311 d	complete cds	2e-62	2506357	dioxygenase	3.4	

F	Nearest	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC		THE TOUR (DIASUT 45. C	Jon Dank)	remest (verginor (Diasex vs. (Non-Redundant Proteins)				
ID	7	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
-								
ĺ	ł	Xenopus laevis						
1445	45060505	notchless (nle)	_		(AF069737) notchless [Xenopus			
1443	AF069737	mRNA, complete cds	2e-62	3687833	lae vis]	le-55		
		Homo sapiens nuclear	ļ		nuclear receptor co-repressor N-			
	i	1 7	ŀ	ŀ	CoR - mouse musculus]			
]	i	receptor co-repressor N-CoR mRNA,			>gi 1583865 prf 2121436A			
1446	AF044209	complete cds	5. 62	212502	thyroid hormone receptor co-			
1	711 044209	complete cus	5e-63	2137603	repressor [Mus musculus]	2e-47		
		Human aryl			1			
1		hydrocarbon receptor		, .	ľ			
	1	nuclear translocator			(45001207)			
1	ļ	(ARNT) mRNA,			(AF001307) aryl hydrocarbon	- 1		
1447	M69238	complete cds.	2e-63	2702210	receptor nuclear translocator;	I		
	1	compiete eus.	26-03	2702319	Arnt [Homo sapiens]	5e-19		
1	İ				KINASE ALPHA	Ì		
-		1	1		REGULATORY CHAIN,	i		
1	,]			LIVER ISOFORM			
1	f]]	1		(PHOSPHORYLASE KINASE	Į.		
]	l j	i		ALPHA L SUBUNIT)			
1	ĺ]			>gi 663010 (X80497)			
1	1				phosphorylase kinase			
1		H.sapiens PHKLA	, .		phosphorylase kinase alpha	ŀ		
1448	X80497	mRNA	2e-63	1170685	subunit [Homo sapiens]	5e-22		
		Homo sapiens			(AF031141) ubiquitin	36-22		
		ubiquitin conjugating			conjugating enzyme [Homo			
1449	AF031141	епгуте	2e-63	2623260	sapiens	1e-23		
				·				
1 1			1		PROBABLE ATP-	1		
} I	İ	1	1		DEPENDENT RNA			
]]	ſ	1			HELICASE P47	-		
ŀ			1	•	>gi 2135840 pir 137201 nuclear			
					RNA helicase (DEAD family)	}		
		H.sapiens BAT1	j		BAT1 - human >gi 587146	1		
1450		mRNA for nuclear			(Z37166) nuclear RNA helicase			
1450	Z37166	RNA helicase Human helix-loop-	6 c -64	2500529	(DEAD family) [Homo sapiens]	9e-24		
		helix zipper protein	i					
1		(max) mRNA,	- 1]	ł		
		complete cds. > ::			1	- 1		
l I		sb 141138 141138	ł		1]		
	4.7	Sequence 1 from	- 1					
1		patent US 5624818 >	1					
1		: gb I77062 I77062	Í		!			
		Sequence 1 from	1	j		ı		
1451		patent US 5693487	50.64		Myc-binding factor Max, short			
		ALC: IL US 307340 /	5e-64	88175	form - human	8e-22		

.5	Nearest	Neighbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ		Troighbot (Blasar Vs. C	T T	IAemest IAeiği	gnoof (Blasta Vs. Non-Redundant Proteins)		
Ð	ACCESSION	DESCRIPTION	PVALUE	ACCESSION	DESCRIPTION	P VALUE	
<u> </u>					PROCOLLAGEN-LYSINE.2-	7	
ł	ł	Homo sapiens lysyl			OXOGLUTARATE 5-	1	
1	i	hydroxylase (partial			DIOXYGENASE		
ļ	1	clone 2.2 Kb LH)	1		PRECURSOR (LYSYL	ł	
		RNA, complete	1		HYDROXYLASE) lysyl	İ	
1452	M98252	mature peptide.	2e-64	400205	hydroxylase [Homo sapiens]	7e-22	
1		Ī			ÓVIDÚCT-SPECIFIC		
	i	1,,			GLYCOPROTEIN]	
ì		Human oviductal			PRECURSOR (OVIDUCTAL	1	
1453	TIOOSEO	glycoprotein mRNA,			GLYCOPROTEIN)	•	
1433	U09550	complete cds.	8e-65	2493676	(OVIDUCTIN)	2e-11	
					resiniteratoxin-binding protein	1	
1			٠.		RBP-26, cytosolic - rat		
1	1				>gi 311660 (X67877) cytosolic	l . i	
1		R.norvegicus mRNA			resiniferatoxin binding protein		
		for cytosolic			RBP-26 [Rattus norvegicus]	[·	
		resiniferatoxin-			>gi 1093373 prf 2103310A	1	
1454	X67877	binding protein	7e-65	400777	resiniferatoxin-binding protein		
1.75	X07877	onding protein	/e-03	423664	[Rattus norvegicus]	2e-40	
		Homo sapiens mRNA					
		for KIAA0711			physomina/alous—is said with		
1455	AB018254	protein, complete cds	6e-65	92298	glutamine/glutamic acid-rich		
		Human 40-kDa	00-03	92296	protein	0.98	
		keratin intermediate	ŀ				
		filament precursor	1		keratin 19, type I, cytoskeletal -		
1456	J03607	gene.	3e-65	1070608	human sapiens]	4e-07	
					Truthan sapiens)	46-07	
1 1		Human gamma-	· I			i	
		glutamyl carboxylase	į			•]	
1457	U65896	gene, complete cds	2e-65	<none></none>	<none></none>	<none></none>	
		1			ISOCITRATE		
1 1			i		DEHYDROGENASE (NAD),		
1	!]		MITOCHONDRIAL SUBUNIT	ĺ	
ł	İ		j		ALPHA PRECURSOR		
	1				(ISOCITRIC		
	_		1		DEHYDROGENASE) (NAD+-		
	1	Human NAD(H)-	I		SPECIFIC ICDH)		
	1.	pecific isocitrate	1		dehydrogenase alpha chain	į	
		lehydrogenase alpha			precursor - human >gi 706839	f	
1463		subunit precursor	1		subunit precursor [Homo		
1458	U07681 r	nRNA, complete cds.	2e-65	1708399	sapiens]	4e-26	
- 1	I	luman zinc tinger	}				
1		protein (LD5-1) gene.]			ł	
1459		xons 4, 5 and 6, and			(U57796) zinc finger protein		
1429	U88080 c	omplete cds	2e-65	1373394	[Homo sapiens] >gi 2306773	2e-39	

				T	•			
		Neighbor (BlastN vs. C	enbank)	Nearest Neig	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC]				
ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
<u> </u>					tensin - chicken (fragment)			
1		Gallus domesticus			>gi 63805 (Z18529) tensin			
1460	10000	tensin mRNA	1		[Gallus gallus] >gi 212755			
1460	M96625	sequence.	3e-66	2134419	(L06662) tensin [Gallus gallus]	le-51		
1		Mus musculus myelin						
1	1	gene expression			(4112262)			
1		factor (MEF-2)			(U13262) myelin gene			
1461	U13262	mRNA, partial cds.	1e-70	536926	expression factor [Mus	1		
		Mus musculus Tera	10-70	330920	musculus]	9e-42		
		(Tera) mRNA.						
1462	U64033	complete cds	5e-72	1575505	(U64033) Tera [Mus musculus]	9e-34		
		M.musculus mRNA			TESTIN 2 (TES2)	90-34		
1463	X78989	for testin	6e-74	1351218	[CONTAINS: TESTIN I	8e-31		
		Mus musculus Tera				3031		
1		(Tera) mRNA,			·			
1464	U64033	complete cds	2e-74	1575505	(U64033) Tera [Mus musculus]	5e-37		
i		C 6						
l		Canis familiaris UDP N-acetylglucosamine						
		transporter mRNA,			(AF057365) UDP N-			
1465	AF057365	complete cds	9e-79	2200605	acetylglucosamine transporter			
- 193	111 03 7 3 0 3	Rattus norvegicus	96-79	3298605	[Canis familiaris]	9e-10		
		mRNA for coronin-	i		(AJ006064) coronin-like protein			
1466	AJ006064	like protein	le-82	3757680	[Rattus norvegicus]	3e-62		
		Macaca fascicularis		0,5,000	[Kukus noi vegreus]	36-02		
		UDP-	ļ					
		glucuronosyltransfera	1		KARYOGAMY PROTEIN			
		se mRNA, complete			KAR4 yeast (Saccharomyces			
1467	U91582	cds	4e-89	140396	cerevisiae)	1e-08		
1468	l l	Mouse Hox2.3			HOMEOBOX PROTEIN HOX-			
1408	X06762	mRNA	3e-92	123255	B7 (HOX-2C)	9e-23		
	į.	Cricetulus griseus	. [•				
I		mRNA for	[1	ŀ		
ŀ		Phosphatidylglycerop	ŀ		(AB016930)	i		
		hosphate synthase,	1		Phosphatidylglycerophosphate	l		
1469		complete cds	5e-94	4159682	synthase [Cricetulus griseus]	7e-3 - ∔		
				7137002	SER/THR-RICH PROTEIN	/e-3 4		
			ł		TIO IN DGCR REGION			
		M.musculus T10	ŀ		>gi 480900 pir S37488 gene			
1470	X74504	nRNA	7e-97	1711658	T10 protein - mouse	3e-59		

1.22	Neares	t Neighbor (BlastN vs.	Genhank)	None N	II. (De las	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEC		Signosi (Diusti 4 Vs.	- Contounk)	ivearest Neig	noor (BlastX vs. Non-Redundant P	roteins)			
iD	-	N DESCRIPTION	1			ł			
-	TACCESSIO	N DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE			
<u> </u>			 		OBIQUITIN-COMOONTING				
	1		1	į.	ENZYME E2-17 KD 3 (UBIQUITIN-PROTEIN				
İ	f		i		LIGASE) (UBIQUITIN]			
I	1		f		CARRIER PROTEIN)	1			
ł	I				(E2(17)KB 3)	ł . l			
1	ł	1	1		>gi 1085588 pir S53358	i i			
1	ł			İ	ubiquitin conjugating enzyme	ł			
	1		1		(E217kB) - rat >gi 595666	i i			
	I	Rattus norvegicus	İ	ł	(U13175) ubiquitin conjugating	1			
1	1	clone ubc 10a	1	i	enzyme [Rattus norvegicus]	. [
Í		ubiquitin conjugating	1		norvegicus] >gi 1145691] `]			
1471	U13175	enzyme (E217kB) mRNA, complete cds	. 3e-98		(U39318) UbcH5C [Homo]]			
	015175	h-lamp-2=lysosome-	3e-98	1351345	sapiens	5e-05			
ŀ		associated membrane			1	l [-			
ł		protein-2 protein-2b			j				
1	1	(LAMP2) mRNA,	l						
1	I .	alternatively spliced	I						
1		form h-lamp-2b,							
1472	S79873	complete cds.	e-119	<none></none>	<none></none>	<none></none>			
1		Dos - DNIA for - 24	}						
1473	D13623	Rat mRNA for p34 protein, complete cds	- 112	1000==	ribosome-binding protein p34 -	1			
-	2.3023	protein, complete cus	e-112	480379	rat sp.]	2e-05			
ł		Mus musculus mRNA				1			
		for 49 kDa zinc finger			(AB013357) 49 kDa zinc finger	1			
1474	AB013357	protein, complete cds	e-136	4153886	protein	5e-08			
1						30 00			
		Cricetulus griseus	ı			f			
		mRNA for Phosphatidylglycerop	J						
		hosphate synthase.	ŀ		(AB016930)				
1475		complete cds	e-117	A1 EOCBO	Phosphatidylglycerophosphate	ſ			
			C-11/	4159682	synthase [Cricetulus griseus]	4c-32			
		Rattus norvegicus	-		TRANSLATION INITIATION				
		initiation factor eIF-	1		FACTOR EIF-2B GAMMA				
		2B gamma subunit	1		SUBUNIT (EIF-2B GDP-GTP				
,,,,,		(eIF-2B gamma)			EXCHANGE FACTOR)	- 1			
1476	U38253	mRNA, complete cds	e-103		subunit [Rattus norvegicus]	3e-42			

	Neare	st Neighbor (BlastN vs.	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SE			T T	Mearest Neig	real est regimber (BlastA vs. Non-Redundant Proteins)			
ID	-	DESCRIPTION	P VALUE	4.000000000				
_	1	DEGERIA HOR	TVALUE	ACCESSION	DESCRIPTION	P VALUE		
				ļ	(H3.3Q) histone H3.3 - fruit fly-	ļ		
	1	}	İ		(Drosophila melanogaster)			
	I			1	histone H3.3B - chicken			
1	1		l .		>gi 2119023 pir S61218 histone			
	1		ŀ		H3.3 - fruit fly (Drosophila			
ł	1			[hydei) 1-136) [Oryctolagus			
i i	ı				cuniculus] >gi 8046 (X53822)			
ı	ı		1		Histone H3.3Q gene product			
1	1				[Drosophila melanogaster]			
j	ł		j i		>gi 51198 gallus] >gi 161190			
			:	•	(M17876) histone H3 [Spisula			
1					solidissima] >gi 211853 (M11393) histone 3.3 [Gallus			
	1]	!	[M11393) histone 3.3 [Gallus gallus] >gi 306848 (M11354)			
1	l				H3.3 histone [Homo sapiens]			
ļ	1				melanogaster] >gi 963031			
1					(X81205) histone H3.3 H3.3A			
1		R.norvegicus mRNA			variant [Drosophila	-		
1477	X73683	for histone H3.3	e-117	122075	melanogaster] musculus]	1e-45		
		Rattus norvegicus	[[rsec8 - rat (fragment)			
1478	U32498	rsec8 mRNA, partial cds	e-108	2142062	>gi 1019441 (U32498) rsec8	_		
		100	C-108	2143962	[Rattus norvegicus]	7e-48		
İ	ŀ	Mus musculus ancient				I		
1		ubiquitous 46 kDa	ł			1		
		protein AUP1	j		(U41736) ancient ubiquitous 46			
1479	7741506	precursor (Aup1)			kDa protein AUP46 precursor			
14/9	U41736	mRNA, complete cds	e-146	1517822	[Mus musculus]	5e-49		
		Bos taurus vacuolar	Ī					
		proton pump subunit	ŀ		1			
	•	SFD alpha isoform			(AF041338) vacuolar proton	1		
		(SFD) mRNA,		•	pump subunit SFD alpha			
1480	AF041338	complete cds	e-119	2895578	isoform [Bos taurus]	3e-49		
				****		-5- /2		
		Mus musculus NSD1	j			- 1		
1481	AF064553	protein mRNA,			(AF064553) NSD1 protein			
2 70 1	1 T OO+233	complete cds	e-121	3329465	[Mus musculus]	2e-50		
		Rattus sp. mRNA for	- 1					
l		CDP-diacylglycerol	i		(U41736) ancient ubiquitous 46	j		
		synthase, complete			kDa protein AUP46 precursor	1		
1482		cds	e-146	1517822	[Mus musculus]	2e-51		
		Mouse mRNA for						
1402		Dhm I protein.	1		mouse Dhm I protein - mouse			
1483	D38517	complete cds	e-118	2137562	musculus]	6e-54		

	Neares	Neighbor (BlastN vs.	Genbank)	Nearest Nai	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ				rearest (ver	ghoof (Blastix Vs. Non-Redundant Pr	roteins)		
B	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE		
1.101	ļ.,	M.domesticus MD6			CDC4 repeat unit-containing			
1484	X54352	mRNA	e-139	1085499	protein - mouse	le-55		
		Mus musculus N- terminal asparagine			PROTEIN N-TERMINAL ASPARAGINE AMIDOHYDROLASE (PROTEIN NH2-TERMINAL ASPARAGINE DEAMIDASE) (NTN-AMIDASE) (PNAD) (PROTEIN NH2-TERMINAL ASPARAGINE AMIDOHYDROLASE) (PNAA) > gi 1373365 (U57691)			
		amidohydrolase	1		N-terminal asparagine			
1485	U57692	(Ntan1) mRNA, complete cds	e-118	2498797	amidohydrolase [Mus musculus]			
1486	X80169	M.musculus mRNA for 200 kD protein	e-119	1717793	amidohydrolase [Mus musculus] PROTEIN TSG24 (MEIOTIC CHECK POINT REGULATOR) >gi 1083553 pir A55117 tsg24	5e-57 9e-58		
1487	U57692	Mus musculus N- terminal asparagine amidohydrolase (Ntan1) mRNA, complete cds	e-120	2498797	PROTEIN N-TERMINAL ASPARAGINE AMIDOHYDROLASE (PROTEIN NH2-TERMINAL ASPARAGINE DEAMIDASE) (NTN-AMIDASE) (PNAD) (PROTEIN NH2-TERMINAL ASPARAGINE AMIDOHYDROLASE) (PNAA) >gi 1373365 (U57691) N-terminal asparagine amidohydrolase [Mus musculus] amidohydrolase [Mus musculus]	8e-58		
488	U08215 r	Mus musculus Hsp70- elated NST-1 (hsr.1) nRNA, complete cds.	e-109	473407	(U08215) NST-1 [Mus	7ė-58		
489	D85926 F	Mouse mRNA for Ray, complete cds	e-110	1944389	(D85926) Ray [Mus musculus]			
490	F d e n	Rattus norvegicus ihydroxypolyprenylb nzoate nethyltransferase nRNA, complete cds	e-123		(L20427) dihydroxypolyprenylbenzoate methyltransferase dihydroxypolyprenylbenzoate methyltransferase [Rattus	2e-58		
	N	1.musculus mRNA	C-123	457372	norvegicus) (X56044) protein Htf9C [Mus	4e-59		
191	X56044 fo	or protein Htf9C	e-121	3183977	musculus]	le-60		

	Nearest	Neighbor (BlastN vs. G	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ							
ID	ACCESSION	ESSION DESCRIPTION		ACCESSION	DESCRIPTION	P VALUE	
<u> </u>					PROTO-ONCOGENE		
					TYROSINE-PROTEIN		
1					KINASE FYN (P59-FYN)	J	
Í :		p59fyn(T)=OKT3-			>gi 420217 pir A44991 protein-	i :	
1		induced calcium		ĺ	tyrosine kinase (EC 2.7.1.112)	1	
1492	<u>\$74774</u>	influx regulator	e-163	729896	fyn - mouse	8e-63	
		Mus musculus BUB2-		ł			
1		like protein I					
		(HBLPI) mRNA,			(U88873) BUB2-like protein I		
1493	U88873	complete cds	e-123	4099611	[Mus musculus]	le-63	
i i		Cricetulus griseus HT				. 1	
		protein mRNA,		• •	(U48852) HT protein		
1494	U48852	complete cds.	e-117	1216486	[Cricetulus griseus]	7e-64	
		Rattus norvegicus					
1406	A E02044	rexo70 mRNA,			(AF032667) rexo70 [Rattus		
1495	_AF032667	complete cds	e-142	2827160	norvegicus]	5e-66	
i I		ļ			PHOSPHATIDYLSERINE DECARBOXYLASE		
					PROENZYME		
		Chinese hamster			J .		
		phosphatidylserine			>gi 109423 pir A38732		
		decarboxylase			phosphatidylserine		
1496	M62722	mRNA, 3' end.	e-114	118910	decarboxylase (EC 4.1.1.65) -	2 (2	
1470	14102722	HIRITA, 5 enu.	6-114	118910	Chinese hamster (fragment)	2e-67	
		Mus musculus fatty			(AF072758) fatty acid transport		
		acid transport protein			protein 3; FATP3 [Mus		
1497	AF072758	3 mRNA, partial cds	e-130	3335567	musculus]	le-67	
		J. Harrist, partial val		3333307	inusculus	16-07	
		Rattus norvegicus	j				
l i		mRNA for atypical	. [(AB005549) atypical PKC		
		PKC specific binding	ı		specific binding protein [Rattus		
1498		protein, complete cds	e-113	3868778	norvegicus]	2e-69	
		Mus musculus					
1	·	homeobox protein					
		Meis3 mRNA,			HOMEOBOX PROTEIN	Ì	
1499	U57344	complete cds	e-143	3024124	MEIS3	6e-72	
		Mus musculus SKD3	1		SKD3 PROTEIN SKD3 [Mus		
1500		mRNA, complete cds.	e-142	2493735	musculus]	1e-72	
- 1		Mus musculus					
]		muskelin mRNA,			(U72194) muskelin [Mus	i	
1501	U72194	complete cds	e-148	3493462	musculus]	2e-74	
ļ					PROTEIN TSG24 (MEIOTIC		
- 1	1.				CHECK POINT		
1500		M.musculus mRNA	[REGULATOR)	,	
1502	X80169	for 200 kD protein	e-155	1717793	>gi 1083553 pir A55117 tsg24	3e-77_	

1.504	Nearest Neighbor (BlastN vs. Genbank) Nearest Neighbor (BlastX vs. Non-Redunda					
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
		Mus musculus				
1503		muskelin mRNA. complete cds	e-154	3493462	(U72194) muskelin [Mus musculus]	2e-78
1504		Cricetulus griseus mRNA for Zn finger factor	e-146	3150148	(Y12836) Zn finger factor [Cricetulus griseus]	3e-83

Table 5

SEQ ID	Start	Stop	Score	Direction	Description
29	295	421	5872	For	mkk like kinases
30	31	182	3943	For	Basic region plus leucine zipper transcription factors
31	298	397	5625	For	mkk like kinases
186	175	395	7660	For	SH2 Domain
187	358	432	4320	For	Ank repeat
196	37	322	6049	For	mkk like kinases
234	23	121	4607	For	SH3 Domain
308	110	172	4150	For	Zinc finger, C2H2 type
410	42	191	4036	For	Basic region plus leucine zipper transcription factors
431	71	428	5538	Rev	ATPases Associated with Various Cellular Activities
552	116	288	3930	Rev	Basic region plus leucine zipper transcription factors
639	157	561	5797	For	ATPases Associated with Various Cellular Activities
746	209	427	5379	For	Fibronectin type III domain
768	116	288	3930	For	Basic region plus leucine zipper transcription factors
807	339	392	3620	For	Zinc finger, C2H2 type
820	341	406	2930	Rev	EF-hand
822	108	262	4179	For	Basic region plus leucine zipper transcription factors
836	158	353	4430	For	Basic region plus leucine zipper transcription factors
1157	41	444	5279	Rev	protein kinase
1192	186	416	5469	For	Fibronectin type III domain
1268	238	315	3540	For	Ank repeat
1269	79	240	11640	For	LIM domain containing proteins
1288	73	234	3953	For	Basic region plus leucine zipper transcription factors

SEQ ID	Start	Stop	Score	Direction	Description
1309	248	404	8226	for	LIM domain containing proteins
1324	294	356	4690	for	Zinc finger, C2H2 type
1325	1	234	8981	for	C2 domain (prot. kinase C like)
1336	66	164	6390	for	WD domain, G-beta repeats
1360	222	377	8686	for	LIM domain containing proteins
1365	69	257	5221	for	Basic region plus leucine zipper transcription factors
1380	42	140	7130	for	WD domain, G-beta repeats
1386	243	398	8736	for	LIM domain containing proteins
1410	222	350	10553	for	Trypsin
1417	8	354	6073	for	Protein Tyrosine Phosphatase
1454	49	209	3996	for	Basic region plus leucine zipper transcription factors
1464	4	180	4978	for	RNA recognition motif. (aka RRM, RBD, or RNP domain)
1478	54	437	5176	for	protein kinase
1496	241	520	3929	for	Helicases conserved C-terminal domain
1496	40	612	5187	for	protein kinase
1503	154	216	4870	for	Zinc finger, C2H2 type
1514	2	252	4662	for	RNA recognition motif. (aka RRM, RBD, or RNP domain)
1527	156	212	3520	for	Zinc finger, C2H2 type
1538	9	635	11087	for	wnt family of developmental signaling proteins
1540	289	471	4107	for	Basic region plus leucine zipper transcription factors
1549	200	391	4118	for	Basic region plus leucine zipper transcription factors
1556	163	354	3958	for	Basic region plus leucine zipper transcription factors
1557	207	398	4038	for	Basic region plus leucinc zipper transcription factors
1563	107	298	3978	for	Basic region plus leucine zipper transcription factors

SEQ ID	Start	Stop	Score	Direction	Description
1622	180	365	4022	for	Basic region plus leucine zipper transcription factors
1630	100	291	3998	for	Basic region plus leucine zipper transcription factors
1674	196	258	4880	for	Zinc finger, C2H2 type
1676	9.	86	6610	for	Homeobox Domain
1677	316	369	5780	rev	Thioredoxins
1688	109	410	17414	for	Ras family
1704	184	372	3977	for	Basic region plus leucine zipper transcription factors
1707	92	439	24100	rev	Phosphatidylinositol-specific phospholipase C, Y domain
1711	263	361	6400	for	WD domain, G-beta repeats
1744	238	433	10572	rev	Serine carboxypeptidases
1755	281	367	2580	for	EF-hand
1762	236	334	5880	for	WD domain, G-beta repeats
1779	64	126	4790	for	Zinc finger, C2H2 type
1801	295	351	4030	for	Zinc finger, C2H2 type
1804	301	378	3460	for	Ank repeat
1808	36	161	4170	for	Basic region plus leucine zipper transcription factors
1811	184	315	8390	for	N-terminal homology in Ets domain
1814	127	294	10770	for	Bromodomain (conserved sequence found in human, Drosophila and yeast proteins.)
1818	9	146	4741	for	Double-stranded RNA binding motif
1819	278	355	3460	for	Ank repeat
1820	123	299	12150	for	Homeobox Domain
1821	127	303	12180	for	Homeobox Domain
1830	184	267	4270	for	Ank repeat
1832	18	173	8987	for	SH3 Domain
1835	51	206	8987	for	SH3 Domain
1839	224	307	4270	for	Ank repeat
1846	12	398	36700	for	G-protein alpha subunit

SEQ ID	Start	Stop	Score	Direction	Description
1909	160	258	6370	for	WD domain, G-beta repeats
1911	35	151	9335	for	Zinc finger, C3HC4 type (RING finger)
1980	60	197	7917	for	Zinc finger, C3HC4 type (RING finger)
2065	253	306	5410	for	Zinc finger, CCHC class
2135	2	401	10596	for	ATPases Associated with Various
2133	-	.01	100,0		Cellular Activities
2216	90	179	5380	for	WW/rsp5/WWP domain containing
22.0	, ,				proteins
2218	127	225	5500	for	WD domain, G-beta repeats
2281	20	387	6044	for	Protein Tyrosine Phosphatase
2282	183	353	5136	for	C2 domain (prot. kinase C like)
2286	12	382	5228	for	protein kinase
2310	20	371	5962	for	Protein Tyrosine Phosphatase
2363	48	211	4132	for	Basic region plus leucine zipper
2505					transcription factors
2424	43	194	3996	for	Basic region plus leucine zipper
					transcription factors
2428	25	350	4675	for	Dual specificity phosphatase, catalytic
					domain
2562	18	101	4560	for	Ank repeat
2577	0	311	10295	for	4 transmembrane segments integral
					membrane proteins
2591	60	165	4560	for	SH2 Domain
2684	9	461	5759	for	ATPases Associated with Various
					Cellular Activities
2826	116	400	16107	for	DEAD and DEAH box helicases
2859	100	320	5550	rev	ATPases Associated with Various
					Cellular Activities
2871	198	392	9384	for	DEAD and DEAH box helicases
2944	18	281	10480	for	Calpain large subunit, domain III
2969	5	387	5976	rev	protein kinase
3015	131	214	3600	for	Ank repeat
3047	191	292	5295	for	WD domain, G-beta repeats
3081	190	252	4360	for	Zinc finger, C2H2 type
3108	275	367	5791	for	WD domain, G-beta repeats
3147	190	369	4022	for	Basic region plus leucine zipper
					transcription factors
3152	129	320	3947	for	Basic region plus leucine zipper
					transcription factors
3158	167	334	4180	for	Basic region plus leucine zipper
					transcription factors
3175	14	164	5951	for	mkk like kinases

SEQ ID	Start	Stop	Score	Direction	Description
3175	8	112	5968	for	protein kinase
3178	45	386	19398	for	ATPases Associated with Various
					Cellular Activities
3183	14	215	9133	for	4 transmembrane segments integral
					membrane proteins
3190	229	390	6089	for	mkk like kinases
3190	118	390	8063	for	protein kinase
3193	293	355	3570	for	Zinc finger, C2H2 type
3195	0	215	10146	for	4 transmembrane segments integral
					membrane proteins
3197	281	343	4490	for	Zinc finger, C2H2 type
3208	34	256	4190	for	Basic region plus leucine zipper
					transcription factors
3258	138	394	9877	for	Ras family
3266	8	139	9328	for	ATPases Associated with Various
					Cellular Activities
3267	97	180	3820	for	Ank repeat
3274	11	187	15442	for	Fork head domain, eukaryotic
					transcription factors
3281	15	182	9681	for	mkk like kinases
3285	16	102	4680	for	EF-hand
3292	208	300	5585	for	WD domain, G-beta repeats
3297	7	153	6100	for	Helicases conserved C-terminal domain
3306	161	223	4900	for	Zinc finger, C2H2 type
3307	43	321	8740	for	SH2 Domain
3339	94	342	14970	for	SH2 Domain
3345	65	271	12512	for	PDZ domain
3351	124	270	6068	for	Phorbol esters/diacylglycerol binding

Example 4 DIFFERENTIAL EXPRESSION OF POLYNUCLEOTIDES OF THE INVENTION: DESCRIPTION OF LIBRARIES AND DETECTION OF DIFFERENTIAL EXPRESSION

The relative expression levels of the polynucleotides of the invention was assessed in several libraries prepared from various sources, including cell lines and patient tissue samples. Table 6 provides a summary of these libraries, including the shortened library name (used hereafter), the mRNA source used to prepare the cDNA library, the abbreviated name of the library that is used in the tables below (in quotes), and the approximate number of clones in the library.

<u>Table 6</u> Description of cDNA Libraries

Library (lib #)	Description	Number of Clones in this Clustering
1	Km12 L4 Human Colon Cell Line, High Metastatic Potential (derived from Km12C) "High Colon"	307133
2	Km12C Human Colon Cell Line, Low Metastatic Potential "Low Colon"	284755
3	MDA-MB-231 Human Breast Cancer Cell Line, High Metastatic Potential; micro-metastases in lung "High Breast"	326937
4	MCF7 Human Breast Cancer Cell, Non Metastatic "Low Breast"	318979
8	MV-522 Human Lung Cancer Cell Line, High Metastatic Potential "High Lung"	223620
9	UCP-3 Human Lung Cancer Cell Line, Low Metastatic Potential "Low Lung"	312503

Library (lib #)	Description	Number of Clones in this Clustering
12	Human microvascular endothelial cells (HMEC) – Untreated PCR (OligodT) cDNA library	41938
13	Human microvascular endothelial cells (HMEC) – Basic fibroblast growth factor (bFGF) treated PCR (OligodT) cDNA library	42100
14	Human microvascular endothelial cells (HMEC) – Vascular endothelial growth factor (VEGF) treated PCR (OligodT) cDNA library	42825
15	Normal Colon – UC#2 Patient PCR (OligodT) cDNA library "Normal Colon Tumor Tissue"	34285
16	Colon Tumor – UC#2 Patient PCR (OligodT) cDNA library "Normal Colon Tumor Tissue"	35625
17	Liver Metastasis from Colon Tumor of UC#2 Patient PCR (OligodT) cDNA library "High Colon Metastasis Tissue"	36984
18	Normal Colon – UC#3 Patient PCR (OligodT) cDNA library "Normal Colon Tumor Tissue"	36216
19	Colon Tumor – UC#3 Patient PCR (OligodT) cDNA library "High Colon Tumor Tissue"	41388
20	Liver Metastasis from Colon Tumor of UC#3 Patient PCR (OligodT) cDNA library "High Colon Metastasis Tissue"	30956
21	G RRpz Human Prostate Cell Line	164801
22	WOca Human Prostate Cancer Cell Line	162088

The KM12L4 and KM12C cell lines are described in Example 1 above. The MDA-MB-231 cell line was originally isolated from pleural effusions (Cailleau, J. Natl. Cancer. Inst. (1974) 53:661), is of high metastatic potential, and forms poorly differentiated adenocarcinoma grade II in nude mice consistent with breast carcinoma.

. The MCF7 cell line was derived from a pleural effusion of a breast adenocarcinoma and is non-metastatic. The MV-522 cell line is derived from a human lung carcinoma and is of high metastatic potential. The UCP-3 cell line is a low metastatic human lung carcinoma cell line; the MV-522 is a high metastatic variant of UCP-3. These cell lines are well-recognized in the art as models for the study of human breast and lung cancer (see, e.g., Chandrasekaran et al., Cancer Res. (1979) 39:870 (MDA-MB-231 and MCF-7); Gastpar et al., J Med Chem (1998) 41:4965 (MDA-MB-231 and MCF-7); Ranson et al., Br J Cancer (1998) 77:1586 (MDA-MB-231 and MCF-7); Kuang et al., Nucleic Acids Res (1998) 26:1116 (MDA-MB-231 and MCF-7); Varki et al., Int J Cancer (1987) 40:46 (UCP-3); Varki et al., Tumour Biol. (1990) 11:327; (MV-522 and UCP-3); Varki et al., Anticancer Res. (1990) 10:637; (MV-522); Kelner et al., Anticancer Res (1995) 15:867 (MV-522); and Zhang et al., Anticancer Drugs (1997) 8:696 (MV522)). The samples of libraries 15-20 are derived from two different patients (UC#2, and UC#3). The bFGF-treated HMEC were prepared by incubation with bFGF at 10ng/ml for 2 hrs; the VEGF-treated HMEC were prepared by incubation with 20ng/ml VEGF for 2 hrs. Following incubation with the respective growth factor, the cells were washed and lysis buffer added for RNA preparation. The GRRpz cell line refers to low passage (3 passages or fewer) human prostate cells, and the WOca cell line refers to low passage (3 passages or fewer) human prostate cancer cells.

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Each of the libraries is composed of a collection of cDNA clones that in turn are representative of the mRNAs expressed in the indicated mRNA source. In order to facilitate the analysis of the millions of sequences in each library, the sequences were assigned to clusters. The concept of "cluster of clones" is derived from a sorting/grouping of cDNA clones based on their hybridization pattern to a panel of roughly 300 7bp oligonucleotide probes (see Drmanac et al., Genomics (1996) 37(1):29). Random cDNA clones from a tissue library are hybridized at moderate stringency to 300 7bp oligonucleotides. Each oligonucleotide has some measure of specific hybridization to that specific clone. The combination of 300 of these measures of hybridization for 300 probes equals the "hybridization signature" for a specific clone. Clones with similar sequence will have similar hybridization signatures. By developing a sorting/grouping algorithm to analyze these signatures, groups of clones in a library can be identified and brought together computationally. These groups of clones are termed "clusters". Depending on the stringency of the selection in the algorithm (similar to the stringency of hybridization in a classic library cDNA screening protocol), the "purity" of each cluster can be controlled. For example, artifacts of clustering may

occur in computational clustering just as artifacts can occur in "wet-lab" screening of a cDNA library with 400 bp cDNA fragments, at even the highest stringency. The stringency used in the implementation of cluster herein provides groups of clones that are in general from the same cDNA or closely related cDNAs. Closely related clones can be a result of different length clones of the same cDNA, closely related clones from highly related gene families, or splice variants of the same cDNA.

Differential expression for a selected cluster was assessed by first determining the number of cDNA clones corresponding to the selected cluster in the first library (Clones in 1st), and the determining the number of cDNA clones corresponding to the selected cluster in the second library (Clones in 2nd). Differential expression of the selected cluster in the first library relative to the second library is expressed as a "ratio" of percent expression between the two libraries. In general, the "ratio" is calculated by: 1) calculating the percent expression of the selected cluster in the first library by dividing the number of clones corresponding to a selected cluster in the first library by the total number of clones analyzed from the first library; 2) calculating the percent expression of the selected cluster in the second library by dividing the number of clones corresponding to a selected cluster in a second library by the total number of clones analyzed from the second library; 3) dividing the calculated percent expression from the first library by the calculated percent expression from the second library. If the "number of clones" corresponding to a selected cluster in a library is zero, the value is set at 1 to aid in calculation. The formula used in calculating the ratio takes into account the "depth" of each of the libraries being compared, i.e., the total number of clones analyzed in each library.

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In general, a polynucleotide is said to be significantly differentially expressed between two samples when the ratio value is greater than at least about 2, preferably greater than at least about 3, more preferably greater than at least about 5, where the ratio value is calculated using the method described above. The significance of differential expression is determined using a z score test (Zar, <u>Biostatistical Analysis</u> Prentice Hall, Inc., USA, "Differences between Proportions," pp 296-298 (1974)).

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EXAMPLE 5 POLYNUCLEOTIDES DIFFERENTIALLY EXPRESSED IN HIGH METASTATIC POTENTIAL BREAST CANCER CELLS VERSUS LOW METASTATIC BREAST CANCER CELLS

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high metastatic potential breast cancer tissue and low metastatic breast cancer cells. Expression of these sequences in breast cancer can be valuable in determining diagnostic, prognostic and/or treatment information. For example, sequences that are highly expressed in the high metastatic potential cells can be indicative of increased expression of genes or regulatory sequences involved in the metastatic process. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant more aggressive treatment. In another example, sequences that display higher expression in the low metastatic potential cells can be associated with genes or regulatory sequences that inhibit metastasis, and thus the expression of these polynucleotides in a sample may warrant a more positive prognosis than the gross pathology would suggest.

The differential expression of these polynucleotides can be used as a diagnostic marker, a prognostic marker, for risk assessment, patient treatment and the like. These polynucleotide sequences can also be used in combination with other known molecular and/or biochemical markers.

The following tables summarize polynucleotides that are differentially expressed between high metastatic potential breast cancer cells and low metastatic potential breast cancer cells.

Table 7

Differentially expressed polynucleotides: Higher expression in high metastatic potential breast cancer (lib3) relative to low metastatic breast cancer cells (lib4)

SEQ ID NOs:	Lib3 clones	Lib4 clones	lib3/lib4
472	64	0	62
1851	6	0	6
1856	8	0	8
1867	6	0	6
1872	6	0	6
1875	12	3	4
1923	89	22	4

SEQ ID NOs:	Lib3 clones	Lib4 clones	lib3/lib4
2118	7	0	7
2119	7	0	7
2135	37	13	3
2190	19	0	19
2193	16	5	3
2232	12	2	6
2239	6	0	6
2338	21	2	10
2378	16	4	4
2394	6	0	6
2395	6	0	6
2490	13	3	4
2505	16	2	8
2540	8	1	8
2542	11	1	11
2607	11	2	5
2640	22	5	4
2674	8	0	8
2679	19	0	19
2684	14	4	3
2707	8	0	8
2724	9	0	. 9
2757	6	0	6
2776	10	0	10
2804	13	2	6
2818	6	0	6
2906	14	0	14
2959	26	. 8	3
2964	17	4	4
2968	6	0	6
2977	22	3	7
2980	13	1	13
3010	6	0	6
3043	10	1	10
3071	33	12	3
3072	9	1	9
3095	19	3	6
3097	11	2	5
3173	12	2	6
3203	8	1	8
3210	27	8	3

SEQ ID NOs:	Lib3 clones	Lib4 clones	lib3/lib4
3212	13	1	13
3284	8	0	8
3288	6	0	6
3331	14	3	5
3335.	13	1	13

Table 8
Differentially expressed polynucleotides: Higher expression in low metastatic breast cancer cells (lib4) relative to high metastatic potential breast cancer (lib3)

SEQ ID NOs:	Lib 3 Clones	Lib 4 Clones	lib4/lib3
402	0	6	6
614	3	21	7
624	0	6	6
626	0	8	8
712	0	9	9
744	0	7	7
1325	2	29	15
1452	2	13	7
1880	0	9	9
1915	0	7	7
1951	0	6	6
1955	8	32	4
2015	0	7	7
2046	0	7	7
2076	1	22	23
2087	0	6	6
2124	0	9	9
2145	0	8	8
2162	0	6	6
2163	0	12	12
2164	5	19	4
2172	2	15	8
2192	5.	16	3
2244	20	43	2
2266	3	18	6
2313	24	56	2
2346	1	13	13

SEQ ID NOs:	Lib 3 Clones	Lib 4 Clones	lib4/lib3
2355	0	10	10
2371	0	6	6
2393	1	17	17
2404	1	21	22
2443	0	6	6
2460	0	11	11
2523	0	6	6
2575	. 1	10	10
2578	0	6	6
2584	1	17	17
2590	0	6	6
2609	1	9	9
2632	5	24	5
2714	5	24	5
2728	0	6	6
2752	1	14	14
2794	4	15	4
2826	0	7	7
2987	5	15	3
3005	1	14	14
3009	20	58	3
3047	4	17	4
3057	2	17	9
3075	2	11	6
3076	0	6	6
3102	0	6	6
3128	15	52	4
3132	15	52	4
3142	0	6	6
3187	22	49	2
3253	23	96	4
3282	19	46	2
3285	20	40	2
3346	0	9	9

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EXAMPLE 6

POLYNUCLEOTIDES DIFFERENTIALLY EXPRESSED IN HIGH METASTATIC POTENTIAL LUNG
CANCER CELLS VERSUS LOW METASTATIC LUNG CANCER CELLS

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high metastatic potential lung cancer cells and low metastatic lung cancer cells. Expression of these sequences in lung cancer tissue can be valuable in determining diagnostic, prognostic and/or treatment information. For example, sequences that are highly expressed in the high metastatic potential cells can be indicative of increased expression of genes or regulatory sequences involved in the metastatic process. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant more aggressive treatment. In another example, sequences that display higher expression in the low metastatic potential cells can be associated with genes or regulatory sequences that inhibit metastasis, and thus the expression of these polynucleotides in a sample may warrant a more positive prognosis than the gross pathology would suggest.

The differential expression of these polynucleotides can be used as a diagnostic marker, a prognostic marker, for risk assessment, patient treatment and the like. These polynucleotide sequences can also be used in combination with other known molecular and/or biochemical markers.

The following tables summarize polynucleotides that are differentially expressed between high metastatic potential lung cancer cells and low metastatic potential lung cancer cells:

Table 9

Differentially expressed polynucleotides: Higher expression in high metastatic potential lung cancer cells (lib8) relative to low metastatic lung cancer cells (lib9)

SEQ ID NO:	Lib8 clones	Lib9 clones	lib8/lib9
. 14	10	0	10
137	5	0	5
151	5	0	7
152	9	0	13
171	6	0	8
200	10	0	14
254	5	0	7
262	5	0	7
271	5	0	7
348	6	1	8
412	5	0	7
507	5	0	7
520	6	0	8
530	5	0	7
588	5	0	7
623	7	0	10
637	7	0	10
660	5	0	7
678	8	0	11
680	5	0	7
700	9	2	6
714	28	13	3
774	11	0	15
812	5	0	7
834	8	2	6
901	11	2	8
1168	5	0	7
1333	6	0	8
1352	5	0	. 7
1524	11	1	15
1706	5	0	7
1752	17	9	3
1768	20	4	7
1769	5	0	7
1780	6	0	8

SEQ ID NO:	Lib8 clones	Lib9 clones	lib8/lib9
1781	40	3	19
1799	6	1	8
1803	6	1	8
1811	16	9	2
1884	6	0	8
1919	8	1	11
1939	6	^ 0	8
1975	43	9	7
2024	12	1	17
2045	8	1	11
2060	20	13	2
2071	16	4	6
2128	5	0	7
2177	10	2	7
2181	44	13	5
2184	11	1	15
° 2185	10	4	3
2283	7	0	10
2311	10	4	3
2314	10	0	14
2393	14	6	3
2398	6	1	8
2460	10	4	3
2514	. 6	0	8
2597	5	0	7
2657	8	2	6
.2669	6	1	8
2670	6	1	8
3047	21	3	10
3050	16	5	4
3092	7	1	10
3140	181	119	2
3157	5	0	7
3187	16	5	. 4
3210	5	0	7
3220	28	4	10
3236	7	1	10
3249	16	0	22
3264	8	2	6
3305	7	0	10
3309	20	0	28

SEQ ID NO:	Lib8 clones	Lib9 clones	lib8/lib9
3318	24	4	8
3330	5	0	7
3331	5	0	7

Table 10

Differentially expressed polynucleotides: Higher expression in low metastatic lung cancer cells (lib 9) relative to high metastatic potential lung cancer cells (lib 8)

SEQ ID NO:	Lib 8 clones	Lib 9 clones	lib 9/lib 8
24	3	20	5
53	0	18	13
64	0	8	6
70	0	11	8
. 105	10	66	5
129	0	16	11
214	1	14	10
233	4	35	6
237	0	13	9
264	0	29	21
329	- 2	17	6
368	1	37	26
370	0	11	8
418	0	8	6
450	0	9	6
461	0	9	6
484	0	26	19
494	0	41	29
517	1	12	9
522	1	11	8
581	1	17	12
614	3	23	5
706	0	11	8
726	5	23	3
806	0	14	10
824	. 0	9	6
836	1	14	10
874	0	12	9
900	5	21	3
1017	2	14	5

SEQ ID NO:	Lib 8 clones	Lib 9 clones	lib 9/lib 8
1144	0	8	6
1154	0	12	9
1166	2	45	16
1170	1	13	9
1302	2	13	5
1326	1	13	9
1327	1	13	9
1367	0	12	9
1377	0	12	9
1437	2	18	6
1442	1	14	10
1466	0	13	9
1476	0	13	9
1495	0	8	. 6
1496	1	13	9
1664	38	253	. 5
1682	1	17	12
1687	0_	9	6
1758	0	8	6
1817	4	18	3 .
1837	3	16	4
1845	3	23	5
1856	2	17	6
1910	1	18	13
2146	2	16	9
2156	0	9	6
2463	0	12	9
2724	10	38	3
2749	403	2000	4
2801	6	25	3
2993	3	18	4
3080	0	10	7
3107	3	23	5
3292	0	20	14
3324	110	548	4

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EXAMPLE 7

POLYNUCLEOTIDES DIFFERENTIALLY EXPRESSED IN HIGH METASTATIC POTENTIAL COLON CANCER CELLS VERSUS LOW METASTATIC COLON CANCER CELLS

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high metastatic potential colon cancer cells and low metastatic colon cancer cells. Expression of these sequences in colon cancer tissue can provide diagnostic, prognostic and/or treatment information. For example, sequences that are highly expressed in the high metastatic potential cells can be indicative of increased expression of genes or regulatory sequences involved in the metastatic process. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant more aggressive treatment. In another example, sequences that display higher expression in the low metastatic potential cells can be associated with genes or regulatory sequences that inhibit metastasis, and thus the expression of these polynucleotides in a sample may warrant a more positive prognosis than the gross pathology would suggest.

The differential expression of these polynucleotides can be used as a diagnostic marker, a prognostic marker, for risk assessment, patient treatment and the like. These polynucleotide sequences can also be used in combination with other known molecular and/or biochemical markers.

The following table summarizes identified polynucleotides with differential expression between high metastatic potential colon cancer cells and low metastatic potential colon cancer cells:

Table 11

25 Differentially expressed polynucleotides: Higher expression in low metastatic colon cancer cells (lib 2) relative to high metastatic potential colon cancer cells (lib 1)

SEQ ID NOs:	Lib 1 clones	Lib 2 clones	lib 2/lib 1
429	0	9	10
1494	0	.8	9
1923	34	114	4
1986	3	12	4
2018	0	9	10
2036	2	10	5
2049	8	25	3
2135	24	87	4

SEQ ID NOs:	Lib 1 clones	Lib 2 clones	lib 2/lib 1
2146	2	16	9
2208	6	27	5
2215	2	11	6
2239	1	10	11
2307	2	12	6
2313	28	62	2
2357	5	14	3 .
2360	3	21	8
2362	0	6	6
2378	3	12	4
2569	3	20	7
2571	0	6	. 6
2588	54	172	3
2592	15	41	3
2611	0	6	6
2636	0	9	10
2641	7	20	3
2650	0	9	10
2662	0	9	10
2674	4	13	4
2682	0	6	6
2702	9	25	3
2704	8	23	3
2715	2	12 (6
2804	9	22	3
2821	13	-29	2
2840	1	8	9
2846	2	15	. 8
2866	0	6	6
2906	0	6	6
2915	44	109	3
2933	0	6	6
2935	5	16	3
2957	1	11	12
2959	3	27	10
2977	16	30	2
2980	12	27	. 2
3000	2	13	7
3009	12	29	3
3115	0	7	8
3156	502	2170	5

SEQ ID NOs:	Lib 1 clones	Lib 2 clones	lib 2/lib 1
3210	2	21	11
3211	0	9	10
3213	0	7	8
3235	2	12	6
3251	2	12	. 6
3296	3	12	4
3335	1	8	9

EXAMPLE 8

POLYNUCLEOTIDES DIFFERENTIALLY EXPRESSED IN HIGH METASTATIC POTENTIAL

COLON CANCER PATIENT TISSUE VERSUS NORMAL PATIENT TISSUE

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A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high metastatic potential colon cancer tissue and normal tissue. Expression of these sequences in colon cancer tissue can provide diagnostic, prognostic and/or treatment information. For example, sequences that are highly expressed in the high metastatic potential cells can be indicative of increased expression of genes or regulatory sequences involved in the advanced disease state which involves processes such as angiogenesis, dedifferentiation, cell replication, and metastasis. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant more aggressive treatment.

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The differential expression of these polynucleotides can be used as a diagnostic marker, a prognostic marker, for risk assessment, patient treatment and the like. These polynucleotide sequences can also be used in combination with other known molecular and/or biochemical markers.

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The following tables summarize polynucleotides that are differentially expressed between high metastatic potential colon cancer tissue and normal colon tissue:

Table 12
Differentially expressed polynucleotides isolated from samples from two patients (patient 2 and patient 3 and): Lower expression in high metastatic potential colon tissue (patient 2:lib 17; patient 3:lib 20) vs. normal colon tissue (patient 2:lib 15; patient 3:lib 18)

SEQ ID NO:	lib 15 clones	lib 17 clones	lib 15/lib 17
69	19	7	3
123	6	0	6
140	-24	8	3
197	6	0	6
198	113	0	121
254	28	9	3
412	28	9	3
512	11	1	12
641	. 17	7	3
642	7	0	8
954	12	3	4
1011	209	16	14
1024	8	0	9
1040	12	3	4
1055	26	7	4
1106	31	15	2
1125	17	0	18
1129	17	0	18
1138	109	0	117
1244	14	1	15
1253	73	0	78
1283	34	7	5
1285	34	7	5
1339	13	4	3
1474	73	0	78
1505	18	3	6
1553	68	6	12
1554	2542	14	195
1605	2542	14	195
1628	6	0	6
1643	142	4	38
1753	12	0	10
1764	13	0	14

SEQ ID NO:	lib 15 clones	lib 17 clones	lib 15/lib 17
SEQ ID NO:	Lib18 Clones	Lib20 Clones	lib18/lib20
105	28	11	2
198	21	0	18
254	9	0	8
412	9	0	8
1011	11	1	9
1138	14	0	12
1253	23	0	20
1643	18	0	15
1764	12	0	10
3156	140	43	3

Table 13

Differentially expressed polynucleotides isolated from samples from two patients (patient 2 and patient 3): Lower expression in normal colon tissue (patient 2:lib 15; patient 3:lib 18)vs. high metastatic potential colon tissue (patient 2:lib 17; patient 3:lib 20).

SEQ ID NO:	Lib 15 Clones	Lib 17 Clones	lib 17/lib 15
321	3	23	7
363	1	9	8
836	21	99	4
859	6	20	3
885	13	28	2
916	13	28	2
981	2	11	. 5
1226	8	70	8
1308	0	8	7
1317	29	84	3
1429	27	127	4
1442	0	9	8
1534	1	12	11
1540	12	. 43	3
1552	0	7	7
1556	1	9	8
1557	1	9	8
1569	2189	5122	2
1571	6	18	3
1576	3	25	8

SEQ ID NO:	Lib 15 Clones	Lib 17 Clones	lib 17/lib 15
1581	4	22	5
1601	25	157	6
1613	9	48	5
1616	15	61	4
1620	2	17	8
1622	4	99	23
1626	6	35	5
1647	4	22	5
1664	4	28	7
1683	2	18	8
1704	3	15	5
1800	0	7	7
2749	23	60	2
2784	4	14	3
2805	1	9	8
- 2976	3	14	4
3128	18	57	3
3129	26	124	4
3146	64	210	3
3150	940	2267	2
3151	2	15	. 7
SEQ ID NO:	lib 18 clones	lib 20 clones	1ib 20/lib 18
865	0	5	6
1569	1	7	8
1580	1	7	8
1590	1	7	8
2790	0	. 5	6

EXAMPLE 9

POLYNUCLEOTIDES DIFFERENTIALLY EXPRESSED IN HIGH COLON TUMOR POTENTIAL PATIENT TISSUE VERSUS METASTASIZED COLON CANCER PATIENT TISSUE

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from colon cancer tissue and cells derived from colon cancer tissue metastases to liver. Expression of these sequences in colon cancer tissue can provide diagnostic, prognostic and/or treatment information associated with the transformation of precancerous tissue to malignant tissue. This information

can be useful in the prevention of achieving the advanced malignant state in these tissues, and can be important in risk assessment for a patient.

The following table summarizes identified polynucleotides with differential expression between high tumor potential colon cancer tissue and cells derived from high metastatic potential colon cancer cells:

Table 14

Differentially expressed polynucleotides:

Greater expression in metastatic colon tumor tissue (lib 20) vs.

colon tumor tissue (lib 19)

SEQ ID NO:	lib 19 clones	lib 20 clones	lib 20/lib 19
937	0	6	8
976	0	5	7
1520	1	8	11
1546	1	11	15
1550	1	11	. 15
1574	1	8	11
1580	0	7	9
1590	0	7	9
1599	8	21	4
1607	158	632	5
1622	1	7	9

Table 15
Greater expression in colon tumor tissue (lib 19) than metastatic colon tissue (lib 20)

SEQ ID NO:	lib 19 clones	lib 20 clones.	lib 19/lib 20
105	64	11	4
1011	53	1	40
1226	18	4	3
1571	8	0	6
1726	15	3	4
1811	17	2	6
2749	47	6	6
3146	19	2	7
3324	20	1	15

EXAMPLE 10

POLYNUCLEOTIDES DIFFERENTIALLY EXPRESSED IN HIGH TUMOR POTENTIAL COLON CANCER PATIENT TISSUE VERSUS NORMAL PATIENT TISSUE

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A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high tumor potential colon cancer tissue and normal tissue. Expression of these sequences in colon cancer tissue can provide diagnostic, prognostic and/or treatment information associated with the prevention of the malignant state in these tissues, and can be important in risk assessment for a patient. For example, sequences that are highly expressed in the potential colon cancer cells are associated with or can be indicative of increased expression of genes or regulatory sequences involved in early tumor progression. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant closer attention or more frequent screening procedures to catch the malignant state as early as possible.

The following tables summarize polynucleotides that are differentially expressed between high metastatic potential colon cancer cells and normal colon cells:

Table 16

Differentially expressed polynucleotides detected in samples from patient (patient 2)

Higher expression in normal colon tissue (patient 2, lib 15)

vs. tumor potential colon tissue (patient 2:lib16)

SEQ ID NO:	lib 15 clones	lib 16 clones	lib 16/lib 15
69	19	7	3
105	116	54	2
140	24	4	6
197	6	0	6
198	113	3	40
254	28	6	5
412	28	6	5
642	7	0	7
830	10	2	5
938	31	13	3
1011	209	37	6
1095	12	3	4
1125	17	0	18

SEQ ID NO:	lib 15 clones	lib 16 clones	lib 16/lib 15
1129	17	0	18
1138	109	1	115
1253	73	1	77
1283	34	13	3
1285	34	13	3
1339	13	3	5
1453	11	3	4
1474	73	1	77
1505	18	6	3
1554	2542	448	6
1605	2542	448	6
1614	36	14	3
1630	24	9	3
1643	142	2	75
1646	39	14	3
1649	24	8	3
1677	19	6	3
1753	13	0	14
1764	13	0	14
1766	177	65	3
1772	24	8	3

Table 17
Differentially expressed polypeptides detected in samples from patient. Lower expression in normal colon tissue (lib 18) than colon tumor tissue (lib 19)

SEQ ID NO:	lib 18 clones	lib 19 clones	lib 19/lib 18
3146	3	19	6
3150	21	228	10
3324	3	20	6

Table 18
Differentially expressed polypeptides detected in samples from patient. Higher expression in normal colon tissue (lib 18) than colon tumor tissue (lib 19)

SEQ ID NO:	lib 18 clones	lib 19 clones	lib 18/1ib 19
198	21 .	2	12
465	6	0	7
489	6	0	7
745	6	0	7
859	11	2	6
976	7	0	8
1011	209	37	6
1045	8	1	9
1138	14	0	16
1253	23	0	26
1392	16	4	5
1474	23	0	26
1589	6	0	7
1591	22	11	2
1607	386	158	3
1643	18	0	21
1753	12	0	14
1764	12	0	14
SEQ ID NO:	lib 18 clones	lib 19 clones	lib 19/lib 18
105	28	64	2
1011	11	53	4
1226	2	18	8
1251	6	19	3
1559	1	9	8
1571	0	8	7
1608	1	9	8
1766	2	13	6
1782	1	9	8
1811	11	17	15

Table 19
Differentially expressed polynucleotides:
Higher expression in colon tumor tissue
(patient 2, lib 16) vs. normal colon tissue (patient 2, lib 15)

SEQ ID NO:	lib 15 clones	lib 16 clones	lib 16/lib 15
7	1	9	9
164	6	19	3
734	4	15	4
836	21	53	2
928	2	11	5
965	2	11	5
987	2	11	5
1026	7	19	3
1044	4	16	4
. 1119	4	16	4
1226	8	46	5
1227	0	9	9
1251	7	95	13
1316	0	6	6
1429	27	81	3
1442	0	9	9
1540	12	28	2
1553	68	590	8
1560	4	24	6
1577	1	10	9
1588	5	20	4
1610	3 .	13	4
1620	2	23	11
1626	6	23	4
1673	2	15	7
2416	0	7	7
2749	23	54	2
2976	3	14	4
3129	26	64	2
3132	18	54	3

EXAMPLE 11

POLYNUCLEOTIDES DIFFERENTIALLY EXPRESSED IN GROWTH FACTOR-STIMULATED HUMAN MICROVASCULAR ENDOTHELIAL CELLS (HMEC) RELATIVE TO UNTREATED HMEC

A number of polynucleotide sequences have been identified that are differentially expressed between human microvascular endothelial cells (HMEC) that have been treated with growth factors relative to untreated HMEC.

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Sequences that are differentially expressed between growth factor-treated HMEC and untreated HMEC can represent sequences encoding gene products involved in angiogenesis, metastasis (cell migration), and other developmental and oncogenic processes. For example, sequences that are more highly expressed in HMEC treated with growth factors (such as bFGF or VEGF) relative to untreated HMEC can serve as markers of cancer cells of higher metastatic potential. Detection of expression of these sequences in colon cancer tissue can provide diagnostic, prognostic and/or treatment information associated with the prevention of achieving the malignant state in these tissues, and can be important in risk assessment for a patient. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant closer attention or more frequent screening procedures to catch the malignant state as early as possible.

The following table summarizes identified polynucleotides with differential expression between growth factor-treated and untreated HMEC.

Table 20
Differentially expressed polynucleotides:
Higher expression in untreated HMEC (lib 12) vs. bFGF treated HMEC (lib 13)

SEQ ID NO:	lib 12 clones	lib 13 clones	lib 12/lib 13
849	6	0	6
1059	6	0	6
1206	12	2	6
3208	12	0	12

Lower expression in untreated HMEC (lib 12) vs. bFGF treated HMEC (lib 13)

2748	3	12	4
3325	0	6	6

<u>Table 21</u>
Differentially expressed polynucleotides:

Higher expression in untreated HMEC (lib 12) VEGF treated HMEC (lib14)

SEQ ID NO:	lib 12 clones	lib 14 clones	lib 12/lib 14
1150	9	0	9

Lower expression in untreated HMEC (lib 12) vs. VEGF treated HMEC (lib14)

3324	22	50	2

EXAMPLE 12

10 POLYNUCLEOTIDES DIFFERENTIALLY EXPRESSED IN NORMAL PROSTATE CELLS RELATIVE TO PROSTATE CANCER CELLS

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from normal prostate cells and prostate cancer cells. Expression of these sequences prostate tissue suspected of being cancerous can provide diagnostic, prognostic and/or treatment information. These polynucleotide sequences can also be used in combination with other known molecular and/or biochemical markers. The following table summarizes identified polynucleotides with differential expression between high metastatic potential colon cancer cells and low metastatic potential colon cancer cells:

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Table 22
Differentially expressed polynucleotides: normal prostate cell line (lib 21)
vs. prostate cancer cell line (lib 22)
Higher in lib 21

SEQ ID NO:	lib 21 clones	lib 22 clones	lib 21/lib 22
53	17	2	8
1754	22	8	3
1801	7	0	7
1845	22	6	4
446	8	0	8
1410	6	0	6
2060	18	6	3
2143	12	3	4
2632	13	1	13
2899	16	2	8

Higher in lib 22

12

3338

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86	2	13	7
93	0	9	9
687	0	9	9
1269	1	. 15	15
1581	25	74	3
1647	25	74	3
1649	12	27	2
1710	5	16	3
1717	5	16	3
1772	12	27	2
1960	0	6	6
2987	0	6	6
3128	13	42	3
3132	13	42	3
3150	263	962	4
3222	0	6	6
3268	0	6	6

EXAMPLE 13

POLYNUCLEOTIDES DIFFERENTIALLY EXPRESSED ACROSS MULTIPLE LIBRARIES

A number of polynucleotide sequences have been identified that are differentially expressed between cancerous cells and normal cells across two or more tissue types tested (*i.e.*, breast, colon, lung, and prostate). Expression of these sequences in a tissue of any origin can provide diagnostic, prognostic and/or treatment information associated with the prevention of achieving the malignant state in these tissues, and can be important in risk assessment for a patient. These polynucleotides can also serve as non-tissue specific markers of, for example, risk of metastasis of a tumor. The following polynucleotides were differentially expressed but without tissue type-specificity in at least two of the breast, colon, lung, and prostate libraries tested: 53, 105, 355, 412, 614, 836, 1442, 1581, 1647, 1649, 1664, 1772, 1782, 1811, 1845, 1856, 1875, 1923, 2060, 2071, 2135, 2146, 2239, 2313, 2378, 2393, 2416, 2460, 2490, 2632, 2674, 2704, 2724, 2749, 2784, 2804, 2959, 2976, 2977, 2980, 2987, 3009, 3047, 3128, 3129, 3132, 3146, 3150, 3156, 3210, 3324, 3331, and 3335.

Those skilled in the art will recognize, or be able to ascertain, using not more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. Such specific embodiments and equivalents are intended to be encompassed by the following claims.

All publications and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference. The citation of any publication is for its disclosure prior to the filing date and should not be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it is readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

Deposit Information:

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The following materials were deposited with the American Type Culture Collection (ATCC); CMCC = Chiron Master Culture Collection:

cDNA Libraries Deposited with ATCC

		ATCC	CMCC
Tube Number	Deposit Date	Accession No.	Accession No.
ES137	May 30, 2000		
ES138	May 30, 2000		
ES139	May 30, 2000		
ES140	May 30, 2000		·
ES141	May 30, 2000		
ES142	May 30, 2000		
ES143	May 30, 2000		
ES144	May 30, 2000		
ES145	May 30, 2000		·
ES146	May 30, 2000		
ES147	May 30, 2000		
ES148	May 30, 2000		
ES149	May 30, 2000		
ES150	May 30, 2000		
ES151	May 30, 2000		
ES152	May 30, 2000		
ES153	May 30, 2000		
ES154	May 30, 2000		·
ES155	May 30, 2000		
ES156	May 30, 2000		
ES157	May 30, 2000		
ES158	May 30, 2000		_
ES159	May 30, 2000		
ES160	May 30, 2000		
ES161	May 30, 2000		
ES162	May 30, 2000		
ES163	May 30, 2000		
ES164	May 30, 2000		
ES165	May 30, 2000		
ES166	May 30, 2000		
ES167	May 30, 2000	,	

Table 23 lists the clones for each deposit, designated as "tube" number.

This deposit is provided merely as convenience to those of skill in the art, and is not an admission that a deposit is required under 35 U.S.C. §112. The sequence of the polynucleotides contained within the deposited material, as well as the amino acid sequence of the polypeptides encoded thereby, are incorporated herein by reference and are controlling in the event of any conflict with the written description of sequences

herein. A license may be required to make, use, or sell the deposited material, and no such license is granted hereby.

Retrieval of Individual Clones from Deposit of Pooled Clones

Where the ATCC deposit is composed of a pool of cDNA clones, the deposit was prepared by first transfecting each of the clones into separate bacterial cells. The clones were then deposited as a pool of equal mixtures in the composite deposit. Particular clones can be obtained from the composite deposit using methods well known in the art. For example, a bacterial cell containing a particular clone can be identified by isolating single colonies, and identifying colonies containing the specific clone through standard colony hybridization techniques, using an oligonucleotide probe or probes designed to specifically hybridize to a sequence of the clone insert (e.g., a probe based upon unmasked sequence of the encoded polynucleotide having the indicated SEQ ID NO). The probe should be designed to have a T_m of approximately 80°C (assuming 2°C for each A or T and 4°C for each G or C). Positive colonies can then be picked, grown in culture, and the recombinant clone isolated. Alternatively, probes designed in this manner can be used to PCR to isolate a nucleic acid molecule from the pooled clones according to methods well known in the art, e.g., by purifying the cDNA from the deposited culture pool, and using the probes in PCR reactions to produce an amplified product having the corresponding desired polynucleotide sequence.

Table 23

Clone Name	置 Tube
M00001351A:B02	ES 137
M00001356A:H11	ES 137
M00001363D:D09	ES 137
M00001395D:H02	ES 137
M00001439C:H06	ES 137
M00001476B:G10	ES 137
M00001582A:E02	ES 137
M00003750D:E06	ES 137
M00003761C:F02	ES 137
M00003770A:E05	ES 137
M00003786A:A11	ES 137
M00003800A:F09	ES 137
M00003816D:E11	ES 137
M00003902A:C03	ES 137
M00003991C:F06	ES 137

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Clone Name	Tube
M00003995B:E03	ES 137
M00004046C:A08	ES 137
M00004105D:D05	ES 137
M00004139B:B10	ES 137
M00004140D:C03	ES 137
M00004144A:H05	ES 137
M00004152A:C12	ES 137
M00004155D:A10	ES 137
M00004168A:G11	ES 137
M00004197B:H10	ES 137
M00004222C:E03	ES 137
M00004234A:E07	ES 137
M00004239B:F11	ES 137
M00004241B:H07	ES 137
M00004264B:A05	ES 137

Clone Name	Tube
M00004278A:F09	ES 137
M00004282D:C11	ES 137
M00004308C:C06	ES 137
M00004340C:C07	ES 137
M00004354D:E05	ES 137
M00004354D.E03	ES 137
	ES 137
M00004372B:F07	
M00004378A:B10	ES 137
M00004393B:E07	ES 137
M00023282A:C02	ES 137
M00023300D:C11	ES 137
M00023316C:G08	ES 137
M00023333D:C12	ES 137
M00023352B:F03	ES 137
M00023352D:H03	ES 137
M00023376B:G04	ES 137
M00023377B:F01	ES 137
M00023398B:D12	ES 137
M00023399C:E10	ES 137
M00026803A:F08	ES 137
M00026843B:D10	ES 137
M00026850D:F09	ES 137
M00026851B:F01	ES 137
M00026856D:F02	ES 137
M00026857D:G12	ES 137
M00026859D:D01	ES 137
M00026860B:C05	ES 137
M00026865B:A06	ES 137
M00026868C:E11	ES 137
M00026878A:F05	ES 137
M00026882D:G09	ES 137
M00026885A:H09	ES 137
M00026901A:G07	ES 137
M00026914A:H10	ES 137
M00026915B:C06	ES 137
M00026918B:D01	ES 137
M00026922C:B02	ES 137
M00026922C:G03	ES 137
M00026926A:E10	ES 137
M00026927D:F02	ES 137
M00026928D:A03	ES 137
M00026935C:B04	ES 137
M00026941D:A04	ES 137
M00026941D.A04 M00026944B:E03	ES 137
M00026944B.E03	ES 137
1V100020940A.F12	123 137

Clone Name	Tübe
M00026980A:D09	ES 137
M00027016A:B06	ES 137
M00027018A:C09	ES 137
M00027021A:G02	ES 137
M00027022D:G11	ES 137
M00027030C:H06	ES 137
M00027035D:C06	ES 137
M00027049B:F05	ES 137
M00027078A:B02	ES 137
M00027080A:B01	ES 137
M00027085C:E11	ES 137
M00027094A:B03	ES 137
M00027103B:A09	ES 137
M00027108C:B03	ES 137
M00027121D:C05	ES 137
M00027135A:B11	ES 137
M00027136C:C09	ES 137
M00027141C:H03	ES 137
M00027159D:F03	ES 137
M00027162B:F05	ES 137
M00027178B:G09	ES 137
M00027179D:E06	ES 138
M00027181D:A05	ES 138
M00027195C:E04	ES 138
M00027198B:B08	ES 138
M00027200A:F02	ES 138
M00027207B:F07	ES 138
M00027212D:E03	ES 138
M00027228D:A01	ES 138
M00027232D:B08	ES 138
M00027233B:C01	ES 138
M00027236A:E04	ES 138
M00027237C:B08	ES 138 ES 138
M00027248A:C02	<u> </u>
M00027256B:H09	ES 138 ES 138
M00027258A:A07	ł
M00027263A:F10	ES 138 ES 138
M00027292D:F10 M00027297A:C04	ES 138
M00027297A:C04 M00027299B:B12	ES 138
M00027299B:B12 M00027301A:G05	ES 138
M00027301A:G03	ES 138
M00027301B:B08	ES 138
M00027314C:D09	ES 138
M00027319D:B11	ES 138
IVIUUU2/324D:CU3	E2 138

No.	OT LESS
Clone Name	Tube
M00027347C:G07	ES 138
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M00027359B:G05	ES 138
M00027366A:F11	ES 138
M00027379C:B07	ES 138
M00027392B:H02	ES 138
M00027396D:G08	ES 138
M00027398C:F07	ES 138
M00027438C:G07	ES 138
M00027462A:D07	ES 138
M00027462B:H07	ES 138
M00027468A:C09	ES 138
M00027475B:E10	ES 138
M00027476A:C09	ES 138
M00027486A:F06	ES 138
M00027520A:C05	ES 138
M00027525B:D06	ES 138
M00027526D:F03	ES 138
M00027528C:B10	ES 138
M00027537C:B01	ES 138
M00027546C:B10	ES 138
M00027591B:C04	ES 138
M00027596A:A10	ES 138
M00027596C:E06	ES 138
M00027602B:C01	ES 138
M00027615A:F10	ES 138
M00027617B:C12	ES 138
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M00027625A:H01	ES 138
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M00027647C:D03	ES 138
M00027652B:F11	ES 138
M00027668C:H12	ES 138
M00027729D:H06	ES 138
M00027733A:A02	ES 138
M00027741B:F09	ES 138
M00027743A:C03	ES 138
M00027801C:C11	ES 138
M00027813C:F01	ES 138
	ES 138
	ES 138
M00027837C:D09	ES 138
M00028120D:F12	ES 138
M00028066C:D07	ES 138
	L

Clone Name	Tube
M00028184D:G10	ES 138
M00028185B:A06	ES 138
M00028196D:A03	ES 138
M00028201B:H12	ES 138
M00028207D:E09	ES 138
M00028210B:D02	ES 138
M00028212C:B08	ES 138
M00028215D:F03	ES 138
M00028220A:B04	ES 138
M00028314D:F05	ES 138
M00028316B:H12	ES 138
M00028354A:B12	ES 138
M00028354D:A03	ES 138
M00028357A:G10	ES 138
M00028362A:G11	ES 138
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M00028770A:D04	ES 138
M00028772C:B09	ES 138
M00028775D:F03	ES 138
M00028777B:G12	ES 138
M00031368A:E10	ES 138
M00031417C:G09	ES 138
M00031419D:C04	ES 138
M00031485D:G02	ES 138
M00032480B:E10	ES 139
M00032492A:C01	ES 139
M00032495B:D02	ES 139
M00032499C:A01	ES 139 ES 139
M00032508B:H03 M00032510D:F12	ES 139
M00032510D:F12	ES 139
M00032510D.G06	ES 139
M00032530D:C02	ES 139
M00032535D:H01	ES 139
M00032539B:C11	ES 139
M00032539B:C11	ES 139
M00032541D:H08	ES 139
M00032545B:H09	ES 139
M00032545D:G05	ES 139
M00032550D:C02	ES 139
M00032551B:G05	ES 139
M00032577A:C04	ES 139
14100032377A.C04	1010/

Clone Name	-Tuber
M00032578A:G06	ES 139
M00032584A:H08	ES 139
M00032592A:H11	ES 139
M00032597C:B01	ES 139
M00032638C:G08	ES 139
M00032638D:A06	ES 139
M00032668D:G12	ES 139
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M00032731C:C07	ES 139
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M00032766C:A04	ES 139
M00032790B:A07	ES 139
M00032793A:F06	ES 139
M00032797B:G02	ES 139
M00032808B:G10	ES 139
M00032811B:D02	ES 139
M00032829B:E06	ES 139
M00032830D:G03	ES 139
M00032831C:G07	ES 139
M00032853D:G12	ES 139
M00032864B:B09	ES 139
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M00032921B:1108	ES 139
M00032939A:C10	ES 139
M00032939B:E07	ES 139
M00032940A:C02	ES 139
M00032942B:B02	ES 139
M00032944B:B02	ES 139
1V1UUUJ2704C.UUJ	100 107

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M00032994A:A08 ES 139 M00032995C:C05 ES 139 M00033007C:E01 ES 139 M00033019B:E10 ES 139 M00033033C:H01 ES 139 M00033034C:A06 ES 139 M00033034C:F02 ES 139 M00033037D:C11 ES 139 M00033074A:C08 ES 139 M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033205A:F03 ES 139 M00033218A:C04 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033248A:A11 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	Clone Name	Tube #
M00032995C:C05 ES 139 M00033007C:E01 ES 139 M00033019B:E10 ES 139 M00033033C:H01 ES 139 M00033034C:A06 ES 139 M00033034C:F02 ES 139 M00033037D:C11 ES 139 M00033074A:C08 ES 139 M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033223B:A05 ES 139 M00033248A:B02 ES 139 M00033248A:B02 ES 139 M0003326D:A11 ES 139 M00033276B:G08 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00032990B:A11	ES 139
M00033007C:E01 ES 139 M00033019B:E10 ES 139 M00033033C:H01 ES 139 M00033034C:A06 ES 139 M00033034C:F02 ES 139 M00033037D:C11 ES 139 M00033074A:C08 ES 139 M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033189D:F08 ES 139 M0003320D:G06 ES 139 M00033204B:A07 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033231D:B09 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M0003326D:A11 ES 139 M00033276B:G08 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00032994A:A08	ES 139
M00033019B:E10 ES 139 M00033033C:H01 ES 139 M00033034C:A06 ES 139 M00033034C:F02 ES 139 M00033037D:C11 ES 139 M00033074A:C08 ES 139 M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M0003320D:G06 ES 139 M00033204B:A07 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033231D:B09 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M0003326D:A11 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00032995C:C05	ES 139
M00033033C:H01 ES 139 M00033034C:A06 ES 139 M00033034C:F02 ES 139 M00033037D:C11 ES 139 M00033074A:C08 ES 139 M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033205A:F03 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033231D:B09 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033007C:E01	ES 139
M00033034C:A06 ES 139 M00033034C:F02 ES 139 M00033037D:C11 ES 139 M00033074A:C08 ES 139 M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033223B:H07 ES 139 M00033224A:A11 ES 139 M00033248A:B02 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033276B:G08 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033019B:E10	ES 139
M00033034C:F02 ES 139 M00033037D:C11 ES 139 M00033074A:C08 ES 139 M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033226A:A11 ES 139 M00033231D:G10 ES 139 M00033248A:B02 ES 139 M00033248A:B02 ES 139 M00033262D:A11 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033033C:H01	ES 139
M00033037D:C11 ES 139 M00033074A:C08 ES 139 M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033217B:H07 ES 139 M00033217B:H07 ES 139 M00033223B:H07 ES 139 M00033221D:B09 ES 139 M00033231D:G10 ES 139 M00033248A:B02 ES 139 M0003326D:A11 ES 139 M0003326B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033034C:A06	ES 139
M00033074A:C08 ES 139 M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033205A:F03 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033221D:B09 ES 139 M00033231D:G10 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M0003326D:A11 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033034C:F02	ES 139
M00033130B:F06 ES 139 M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033205A:F03 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033223D:B09 ES 139 M00033231D:B09 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033262D:A11 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033037D:C11	ES 139
M00033140D:F06 ES 139 M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033205A:F03 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033226A:A11 ES 139 M00033231D:B09 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033074A:C08	ES 139
M00033173D:C01 ES 139 M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033205A:F03 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033226A:A11 ES 139 M00033231D:G10 ES 139 M00033248A:B05 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033130B:F06	ES 139
M00033176B:E12 ES 139 M00033186C:D11 ES 139 M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033205A:F03 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033226A:A11 ES 139 M00033231D:B09 ES 139 M00033243B:A05 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033140D:F06	ES 139
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M00033189D:F08 ES 139 M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033205A:F03 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033226A:A11 ES 139 M00033231D:B09 ES 139 M00033243B:A05 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033176B:E12	ES 139
M00033202D:G06 ES 139 M00033204B:A07 ES 139 M00033205A:F03 ES 139 M00033217B:H07 ES 139 M00033218A:C04 ES 139 M00033223B:H07 ES 139 M00033223B:H07 ES 139 M00033221D:B09 ES 139 M00033231D:G10 ES 139 M00033243B:A05 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033186C:D11	ES 139
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M00033223B:H07 ES 139 M00033226A:A11 ES 139 M00033231D:B09 ES 139 M00033231D:G10 ES 139 M00033243B:A05 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M0003326D:A11 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033217B:H07	ES 139
M00033226A:A11 ES 139 M00033231D:B09 ES 139 M00033231D:G10 ES 139 M00033243B:A05 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033262D:A11 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033218A:C04	ES 139
M00033231D:B09 ES 139 M00033231D:G10 ES 139 M00033243B:A05 ES 139 M00033246C:E08 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033262D:A11 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033223B:H07	<u> </u>
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M00033246C:E08 ES 139 M00033248A:B02 ES 139 M00033261C:D12 ES 139 M00033262D:A11 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033231D:G10	
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M00033262D:A11 ES 139 M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139		
M00033263B:G04 ES 139 M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033261C:D12	\$
M00033276B:G08 ES 139 M00033185C:D01 ES 139	M00033262D:A11	
M00033185C:D01 ES 139	M00033263B:G04	
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	M00033288B:D12	ES 140
M00033300D:H12 ES 140	M00033300D:H12	
M00033306D:G08 ES 140		L
M00033306D:H09 ES 140		3
M00033308B:G05 ES 140		
M00033343C:H08 ES 140		1
M00033345D:A09 ES 140		
M00033346C:A05 ES 140		
M00033347C:F02 ES 140		
M00033349D:F05 ES 140		
M00033358A:H12 ES 140	M00033358A:H12	
M00033362C:C05 ES 140	M00033362C:C05	ES 140

Clone Name Tube M00033375A:G04 ES 140	
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M00033376A:C12 ES 140	
M00033377D:A05 ES 140	
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M00033432B:H10 ES 140	
M00033437C:A07 ES 140	•
M00033437C:C03 ES 140	
M00033442A:D06 ES 140	_
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M00033446D:B02 ES 140	
M00033446D:B02 ES 140 M00033450C:A02 ES 140	
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M00033457D:A05 ES 140	_
M00033560D:G07 ES 140	
M00033561C:A02 ES 140	_
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M00038259B:G08 ES 140	
M00038259C:H09 ES 140	
M00038272A:G01 ES 140	_
M00038272D:F11 ES 140	-
M00038272D:111 ES 140	
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M00038303C:D02 ES 140	_
M00038303D:E07 ES 140	

Clone Name	Tube
TEE	ES 140
M00038315C:G11 M00038325D:F12	ES 140
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M00038326B:D04	ES 140
M00038327A:C11	
M00038327D:A05	ES 140 ES 140
M00038328D:A03	ES 140
M00038329A:E08 M00038387B:A07	ES 140
M00038514C:H11	ES 140
M00038614C.H11	ES 140
M00038615A:1112	ES 140
M00038618C:C08	ES 140
M00038619B:A03	ES 140
M00038620B:E09	ES 140
M00038620B.E09	ES 140
M00038631D:B02	ES 140
M00038631D:B02	ES 140
M00038633A:D07	ES 140
M00038633A:D07	ES 140
M00038635A:G09	ES 140
M00038635B:C08	E\$ 140
M00038638D:H03	ES 140
M00038639B:C03	ES 140
M00038639D:F07	ES 140
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M00038662B:A12	ES 140
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M00038663D:H10	ES 140
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M00038994A:A10	ES 140
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M00038999D:C11	ES 140
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M00039004B:A06	ES 140
M00039004B:C11	ES 140
M00039005C:H01	ES 141
M00039006D:B01	ES 141
M00039011D:C10	ES 141
M00039013A:C09	ES 141
M00039013D:F02	ES 141
M00039014A:H10	ES 141
M00039014B:C04	ES 141

Clone Name	Tube
M00039015A:D07	ES 141
M00039015B:G10	ES 141
M00039015B:H09	ES 141
M00039015D:H04	ES 141
M00039016A:A02	ES 141
M00039016D:G06	ES 141
M00039024B:B10	ES 141
M00039025A:H09	ES 141
M00039026D:F05	ES 141
M00039028C:B11	ES 141
M00039030B:E02	ES 141
M00039036C:B05	ES 141
M00039039B:E03	ES 141
M00039039B:F09	ES 141
M00039042B:B02	ES 141
M00039043B:E01	ES 141
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M00039050A:H10	ES 141
M00039052C:F07	ES 141
M00039058A:A04	ES 141
M00039058C:H02	ES 141
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M00039061B:F08	ES 141
M00039063B:D08	ES 141
M00039064D:H09	ES 141
M00039066D:G08	ES 141
M00039068B:B04	ES 141
M00039068C:E06	ES 141
M00039070D:C02	ES 141
M00039072C:C03	ES 141
M00039072C:E02	ES 141
M00039079A:A05	ES 141
M00039080C:H06	ES 141
M00039081B:G06	ES 141
M00039082B:A05	ES 141
M00039084C:G07	ES 141
M00039084C:H03	ES 141
M00039084C:H04	ES 141
M00039084D:D07	ES 141
M00039096A:A05	ES 141
M00039096A:E07	ES 141
M00039097D:D06	ES 141
M00039099A:H08	ES 141
M00039104D:C09	ES 141
M00039105C:B08	ES 141
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M00039107C:E04	ES 141
M00039108D:B06	ES 141
M00039112B:C05	ES 141
M00039118B:C05	ES 141
M00039118D:A06	ES 141
M00039120C:C09	ES 141
M00039120C:H03	ES 141
M00039123A:B10	ES 141
M00039124C:F03	ES 141
M00039124C:H02	ES 141
M00039124C:H08	ES 141
M00039126D:A08	ES 141
M00039127A:G11	ES 141
M00039127D:E10	ES 141
M00039129C:D04	ES 141
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M00039135D:F05	ES 141
M00039135D:G02	ES 141
M00039135D:H02	ES 141
M00039139A:C09	ES 141
M00039139C:G12	ES 141
M00039140A:B08	ES 141
M00039140D:A04	ES 141
M00039140D:D09	ES 141
M00039141C:E01	ES 141
M00039142D:B11	ES 141
M00039144C:E06	ES 141
M00039147A:F10	ES 141
M00039156A:B11 M00039158B:G12	ES 141 ES 141
M00039166B:G06	ES 141
M00039167B:H09	ES 141
M00039167B:1103	ES 141
M00039169A:E12	ES 141
M00039170A:B10	ES 141
M00039170C:F05	ES 141
M00039171B:D11	ES 141
M00039177B:D03	ES 141
M00039179A:G09	ES 141
M00039180A:A07	ES 141
M00039196B:H06	ES 141
M00039196D:A07	ES 141
M00039200A:C10	ES 141
M00039211A:C12	ES 141
M00039212C:C12	ES 142

Clone Name	*Rube
M00039213A:D01	ES 142
M00039213B:F05	ES 142
M00039218A:F03	ES 142
M00039218A:103	ES 142
	ES 142
M00039224A:E12	
M00039228A:B05	ES 142
M00039230A:A10	ES 142
M00039230D:D09	ES 142
M00039230D:G12	ES 142
M00039233A:A03	ES 142
M00039238A:B12	ES 142
M00039238D:A08	ES 142
M00039241A:E11	ES 142
M00039249A:C12	ES 142
M00039249C:G11	ES 142
M00039255C:E12	ES 142
M00039257D:C03	ES 142
M00039258B:E06	ES 142
M00039258D:B08	ES 142
M00039260C:G03	ES 142
M00039263D:A12	ES 142
M00039266A:B02	ES 142
M00039266D:F12	ES 142
M00039266D:H04	ES 142
M00039273B:F02	ES 142
M00039273D:B02	ES 142
M00039274B:G07	ES 142
M00039276B:H09	ES 142
M00039277D:G10	ES 142
M00039279B:C11	ES 142
M00039279B:H02	ES 142
M00039279C:B08	ES 142
M00039281D:B04	ES 142
M00039281D:B04	ES 142
M00039284D:B12	ES 142
M00039280A:C00	ES 142
M00039287C.A00	ES 142
M00039288C:B11	ES 142
M00039293B:C11	,
M00039295B:D03	ES 142
M00039297C:H08	ES 142
M00039298B:B06	ES 142
M00039298B:D03	ES 142
M00039298D:B04	ES 142
M00039299B:G12	ES 142

	Property Services (1)
Clone Name	Tube*
M00039300C:C09	ES 142
M00039300C:G04	ES 142
M00039301B:F06	ES 142
M00039303C:F11	ES 142
M00039304D:B09	ES 142
M00039308B:G08	ES 142
M00039310A:C07	ES 142
M00039313D:G04	ES 142
M00039316A:C01	ES 142
M00039318B:B09	ES 142
M00039319B:H12	ES 142
M00039319C:A04	ES 142
M00039322A:F04	ES 142
M00039328D:D07	ES 142
M00039329A:C01	ES 142
M00039329C:B10	ES 142
M00039333D:D09	ES 142
M00039334B:E03	ES 142
M00039335A:E08	ES 142
M00039339A:H07	ES 142
M00039339C:F03	ES 142
M00039340A:D05	ES 142
M00039340B:E07	ES 142
M00039340B:G08	ES 142
M00039341C:H11	ES 142
M00039341D:D07	ES 142
M00039343B:F12	ES 142
M00039344B:G07	ES 142
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M00039345C:C12	ES 142
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M00039381D:C02	ES 142
M00039384C:E02	ES 142
M00039384C:F08	ES 142
M00039385B:E09	ES 142
M00039391D:F08	ES 142
M00039396D:B04	ES 142
M00039397B:H09	ES 142
M00039398A:B10	ES 142
M00039401B:D02	ES 142
M00039402B:E03	ES 142
M00039403A:G12	ES 142
M00039404B:A05	ES 142
M00039407B:G02	ES 142
M00039411C:E07	ES 142
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Előné Name ** *	Libe >
M00039412D:G06	ES 142
M00039414D:G03	ES 142
M00039415D:E01	ES 142
M00039417A:D03	ES 142
M00039417A:E12	ES 142
M00039417A:B12	ES 143
M00039417B:F01	ES 143
	ES 143
M00039417C:G01	ES 143
M00039418B:D08	
M00039420D:D03	ES 143
M00039422D:F04	ES 143
M00039425C:G01	ES 143
M00039425D:E12	ES 143
M00039428C:E01	ES 143
M00039430B:F12	ES 143
M00039431B:F04	ES 143
M00039432C:A01	ES 143
M00039444C:H02	ES 143
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M00039454B:A11	ES 143
M00039455D:H04	ES 143
M00039456A:C08	ES 143
M00039458B:H11	ES 143
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M00039465A:A08	ES 143
M00039472C:B08	ES 143
M00039475C:E10	ES 143
M00039476B:A02	ES 143
M00039477A:B03	ES 143
M00039477A:D03	ES 143
M00039477D:A10	ES 143
M00039611D:D11	ES 143
	ES 143
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M00039616A:B10	ES 143
M00039616B:C01	
M00039619B:D02	ES 143
M00039631A:C10	ES 143
M00039633D:D05	ES 143
M00039636C:D11	ES 143
M00039637C:A10	ES 143
M00039652B:D05	ES 143
M00039655B:H09	ES 143
M00039655C:C07	ES 143
M00039655C:E08	ES 143
M00039660C:C10	ES 143
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A. Clone: Name	Tube
M00039663C:G09	ES 143
M00039664D:G07	ES 143
M00039672D:D10	ES 143
M00039673A:F09	ES 143
M00039675D:B03	ES 143
M00039675D:H05	ES 143
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M00039777C:E05	ES 143
M00039778B:G03	ES 143
M00039778C:A04	ES 143
M00039781D:D10	ES 143
M00039782A:H10	ES 143
M00039785D:G05	ES 143
M00039788A:E03	ES 143
M00039788B:A06	ES 143
M00039788C:A01	ES 143
M00039790B:D03	ES 143
M00039792A:B04	ES 143
M00039793D:C05	ES 143

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Clone Name	
M00039794A:E04	ES 143
M00039795B:H10	ES 143
M00039795D:E10	ES 143
M00039795D:G06	ES 143
M00039797C:G05	ES 143
M00039798B:B02	ES 143
M00039799A:D10	ES 143
M00039801A:H11	ES 143
M00039807A:D01	ES 143
M00039808D:H02	ES 143
M00039810A:H10	ES 143
M00039813B:B01	ES 144
M00039813B:D11	ES 144
M00039815C:F09	ES 144
M00039816B:D04	ES 144
M00039816C:D05	ES 144
M00039820A:F11	ES 144
M00039820A:H11	ES 144
M00039820B:B06	ES 144
M00039827B:F07	ES 144
M00039828B:C05	ES 144
M00039832A:B12	ES 144
M00039835A:F07	ES 144
M00039838A:F05	ES 144
M00039839B:B01	ES 144
M00039839C:E05	ES 144
M00039847A:F06	ES 144
M00039851B:G11	ES 144
M00039851C:D12	ES 144
M00039854B:F09	ES 144
M00039855C:F01	ES 144
M00039857B:G10	ES 144
M00039859A:F06	ES 144
M00039859C:G10	ES 144
M00039864A:A07	ES 144
	ES 144
M00039869B:F06	ES 144
M00039875D:A10	ES 144
M00039876D:H09	ES 144
M00039877C:C03	ES 144
M00039879C:F05	ES 144
M00039879D:B11	ES 144
M00039880A:H11	ES 144
M00039884A:H11	ES 144
	ES 144
1-10003 7003 C.DO1	

Clone Name 24	Tuber
M00039887C:E07	ES 144
M00039887D:C04	ES 144
M00039888B:D03	ES 144
M00039890A:H05	ES 144
M00039894C:H07	ES 144
M00039896C:H01	ES 144
M00039897D:C10	ES 144
M00039898A:A08	ES 144
M00039898D:C06	ES 144
M00039903A:H07	ES 144
M00039903C:D01	ES 144
M00039903C:F03	ES 144
M00039909C:G05	ES 144
M00039909D:C02	ES 144
M00039910C:G10	ES 144
M00039914D:G12	ES 144
M00039915D:C11	ES 144
M00039927A:F04	ES 144
M00039928B:G05	ES 144
M00039936C:C05	ES 144
M00039938C:A08	ES 144
M00039938C:E11	ES 144
M00039940A:D07	ES 144
M00039940D:G08	ES 144
M00039973D:C08	ES 144
M00039973D:D12	ES 144
M00039975C:C11	ES 144
M00039976D:A12	ES 144
M00039978A:G03	ES 144
M00039981A:E08	ES 144
M00039982C:H04	ES 144
M00039983D:A06	ES 144
M00039984A:C02	ES 144
M00039984B:G12	ES 144
M00039984D:G12	ES 144
M00039987A:F09	ES 144
M00039987C:E12	ES 144
M00039987C:G08	ES 144
M00039988A:E06	ES 144
M00039990C:D10	ES 144
M00040004D:B03	ES 144
M00040005B:C11	ES 144
M00040005D:B07	ES 144
M00040007D:A06	ES 144
M00040009D:B07	ES 144
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Clone Name	Paljubel
M00040010A:F10	ES 144
M00040014B:D01	ES 144
M00040014D:D10	ES 144
M00040014D:F03	ES 144
M00040015C:F08	ES 144
M00040016C:H12	ES 144
M00040017A:C06	ES 144
M00040017D:G03	ES 144
M00040019A:E01	ES 144
M00040021A:F09	ES 144
M00040022C:D06	ES 144
M00040026B:F06	ES 144
M00040029A:B03	ES 144
M00040029A:G04	ES 144
M00040031A:E06	ES 144
M00040032A:B03	ES 144
M00040032A:D09	ES 144
M00040037A:E11	ES 145
M00040038D:G04	ES 145
M00040039D:D06	ES 145
M00040040A:A06	ES 145
M00040041C:C09	ES 145
M00040042B:A10	ES 145
M00040047C:F05	ES 145
M00040052D:F12	ES 145
M00040055D:A06	ES 145
M00040055D:B01	ES 145
M00040060C:H10	ES 145
M00040062B:B05	ES 145
M00040070B:B07	ES 145
M00040071B:A10	ES 145
M00040072C:G09	ES 145
M00040076C:D06	ES 145
M00040077D:C11	ES 145
M00040080C:C06	ES 145
M00040081C:E01	ES 145
M00040085D:A10	ES 145
M00040085D:E04	ES 145
M00040087D:F08	ES 145
M00040088C:E10	ES 145
M00040089A:G08	ES 145
M00040089B:E04	ES 145
M00040089C:E06	ES 145
M00040090B:G09	ES 145
M00040092B:F05	ES 145
	A

	Tillibe M
M00040093B:C02	ES 145
M00040093D:D03	ES 145
M00040097A:C12	ES 145
M00040098C:B01	ES 145
M00040098D:E04	ES 145
M00040098D:G12	ES 145
M00040100C:E05	ES 145
M00040100D:B06	ES 145
M00040103B:H10	ES 145
M00040105C:F11	ES 145
M00040106B:B09	ES 145
M00040107B:H07	ES 145
M00040111C:D05	ES 145
M00040115B:A04	ES 145
M00040115B:H12	ES 145
M00040118D:G10	ES 145
M00040121B:C05	ES 145
M00040122D:A02	ES 145
M00040123A:A09	ES 145
M00040124D:H01	ES 145
M00040129D:E10	ES 145
M00040302C:A04	ES 145
M00040304B:F06	ES 145 ES 145
M00040305A:D11 M00040305C:H06	ES 145
M00040307B:F01	ES 145
M00040307E:F10	ES 145
M00040307C:110	ES 145
M00004825D:D05	ES 145
M00004823D:H02	ES 145
M00004839C:H02	ES 145
M00005018A:B05	ES 145
M00005297D:H08	ES 145
M00005308A:D06	ES 145
M00005351C:G05	ES 145
M00005352C:A02	ES 145
M00005358B:B06	ES 145
M00005359A:D04	ES 145
M00005379A:E04	ES 145
M00005382B:F08	ES 145
M00005384A:C11	ES 145
M00005402B:F08	ES 145
M00005445D:B01	ES 145
M00005449B:B10	ES 145
M00005449B:D01	ES 145 ES 145 ES 145 ES 145 ES 145 ES 145 ES 145 ES 145
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Clone Name	* Tube
M00005457C:A03	ES 145
M00005458A:F11	ES 145
M00005498A:H06	ES 145
M00005531D:F06	ES 145
M00005539D:G01	ES 145
M00005555A:A10	ES 145
M00005556B:D02	ES 145
M00005601D:D08	ES 145
M00005614B:B01	ES 145
M00005623A:G02	ES 145
M00005623D:G12	ES 145
M00005625A:C02	ES 145
M00005673B:B12	ES 145
M00005778B:F09	ES 145
M00005805D:D12	ES 145
M00005820C:E04	ES 145
M00006581D:F08	ES 145
M00006599D:B02	ES 145
M00006657C:G05	ES 145
M00006680B:D02	ES 145
M00006712C:H09	ES 145
M00006809B:B09	ES 145
M00006861B:F09	ES 145
M00006866A:D07	ES 146
M00006886D:H02	ES 146
M00006893C:E07	ES 146
M00006897A:H02	ES 146
M00006928D:D07	ES 146
M00006935C:F06	ES 146
M00006968A:G08	ES 146
M00006977C:G04	ES 146
M00006977D:A03	ES 146 ES 146
M00007012D:H08	ES 146
M00007013A:D09	ES 146
M00007026B:H09	ES 146
M00007108B:A02 M00007112C:B10	ES 146
M00007112C:B10 M00007116C:G02	ES 146
M00007116C:G02 M00007124D:H10	ES 146
M00007124D:H10	ES 146
M00007149A:G02	ES 146
M00007157C:F11	ES 146
M00007165B:G11	ES 146
M00007194A:B09	ES 146
M00007194A:B09	ES 146
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Provesor Separate Const.	Muber
Sectione Name *	A STATE OF STREET, ST.
M00007941D:C09	ES 146
M00007943D:C09	ES 146
M00007972B:H12	ES 146
M00007976A:C10	ES 146
M00007992C:F06	ES 146
M00007994A:G02	ES 146
M00008006B:B03	ES 146
M00008026B:C11	ES 146
M00008045A:H02	ES 146
M00008053A:F10	ES 146
M00008063B:A06	ES 146
M00021665B:F12	ES 146
M00021671D:F12	ES 146
M00021852D:A05	ES 146
M00021866D:A03	ES 146
M00021908D:G12	ES 146
M00021919C:A10	ES 146
M00021923C:D11	ES 146
M00021955A:H02	ES 146
M00021964C:E10	ES 146
M00021972D:C11	ES 146
M00022005C:C06	ES 146
M00022015B:B07	ES 146
M00022054A:H03	ES 146
M00022084D:B01	ES 146
M00022099B:D06	ES 146
M00022105C:C12	ES 146
M00022127C:H03	ES 146
M00022135C:B05	ES 146
M00022138A:E05	ES 146
M00022175D:D12	ES 146
M00022178B:D06	ES 146
M00022181C:D01	ES 146
M00022183B:C02	ES 146
M00022184C:C11	ES 146
M00022233C:A12	ES 146
M00022234C:D06	ES 146
M00022247A:E02	ES 146
M00022257A:B09	ES 146
M00022262D:G03	ES 146
M00022264B:G10	ES 146
M00022363C:G12	ES 146
M00022365D:A03	ES 146
M00022373A:B05	ES 146
M00022373C:B07	ES 146
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TO THE RESERVE WAS AND ASSESSED.	SERVICE SERVICE
Clone Name	
M00022391B:E01	ES 146
M00022391D:F10	ES 146
M00022416A:A07	ES 146
M00022421B:C11	ES 146
M00022433A:E02	ES 146
M00022434D:D06	ES 146
	ES 146
M00022444D:G01	ES 146
M00022467C:B12	ES 146
M00022489C:G04	ES 146
M00022492C:A02	ES 146
M00022495D:H08	ES 146
M00022493B:1100 M00022496B:E12	ES 146
M00022490B.E12 M00022499A:B02	ES 146
M00022499A.B02 M00022533A:A08	ES 146
M00022533A:A08 M00022579C:C11	ES 146
	ES 146
M00022597D:A06	
M00022602A:E09	ES 146
M00022615D:G05	ES 146
M00022634D:C08	ES 146
M00022640C:C12	ES 146
M00022641C:H05	ES 146
M00022646A:H10	ES 146
M00022662D:G11	ES 146
M00022667D:B02	ES 146
M00022668B:B12	ES 146
M00022670D:H11	ES 146
M00022671B:A08	ES 146
M00022684A:C02	ES 146
M00022731A:D02	ES 147
M00022739A:B03	ES 147
M00022747D:E03	ES 147
M00022767B:G11	ES 147
M00022785C:G06	ES 147
M00022793D:B01	ES 147
M00022795B:G06	ES 147
M00022797B:G08	ES 147
M00022817A:H02	ES 147
M00022821C:C09	ES 147
M00022823C:C01	ES 147
M00022830D:D01	ES 147
M00022834B:G11	ES 147
M00022854A:B03	ES 147
M00022856C:A07	ES 147
M00022860C:G04	ES 147
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Clone Name	Fa mibe
M00022885C:H05	ES 147
M00022895A:H08	ES 147
M00022910A:A06	ES 147
M00022925C:A08	ES 147
M00022928B:C01	ES 147
M00022930C:E02	ES 147
M00022938B:F07	ES 147
M00022968B:E02	ES 147
M00022976C:F04	ES 147
M00022979A:D05	ES 147
M00022986D:H09	ES 147
M00022997A:F06	ES 147
M00023001C:C08	ES 147
M00023003C:D07	ES 147
M00023007A:H04	ES 147
M00023007C:E10	ES 147
M00023020C:G08	ES 147
M00023024D:F12	ES 147
M00023032A:B05	ES 147
M00023039D:B05	ES 147
M00023042D:D02	ES 147
M00023044B:D02	ES 147
M00023094A:B11	ES 147
M00023100A:E12	ES 147 ES 147
M00039181D:E05	ES 147
M00039184A:D03	ES 147
M00039184B:B09 M00039361B:E01	ES 147
M00039361B.E01	ES 147
M00039365A:C03	ES 147
M00039367B:H02	ES 147
M00039371B:H06	ES 147
M00039371B:1100 M00039372C:D12	ES 147
M00039374B:B07	ES 147
M00039374C:H12	ES 147
M00039374C:H02	ES 147
M00039374C:H02	ES 147
M00039377D:E12	ES 147
M00039377D:E12	ES 147
M00039379A:B03	ES 147
M00039380C:C09	ES 147
M00039482B:G02	ES 147
M00039493A:C04	ES 147
M00039496B:D08	ES 147
M00039496B:H09	ES 147

Clone Name	Tube
M00039497C:C06	ES 147
M00039499C:A04	ES 147
M00039500C:C04	ES 147
M00039505C:E03	ES 147
M00039508A:C12	ES 147
M00039508C:G01	ES 147
M00039510C:G02	ES 147
M00039510C:G02	ES 147
M00039515A:A06	ES 147
	ES 147
M00039515D:C11	ES 147
M00039517B:G12	l
M00039521A:A02	ES 147
M00039521D:H03	ES 147
M00039528B:B12	ES 147
M00039529C:D07	ES 147
M00039530B:E02	ES 147
M00039533A:C12	ES 147
M00039533B:G08	ES 147
M00039533D:F04	ES 147
M00039535D:D10	ES 147
	ES 147
M00039536C:H11	ES 147
M00039561A:B07	ES 147
M00039561B:A09	ES 147
	ES 147
M00039564B:C01	ES 147
	ES 147
	ES 147
M00039584C:C01	ES 147
M00039584C:C11	ES 147
M00039587C:F12	ES 147
M00039590D:D02	ES 147
M00039591C:D06	ES 147
M00039595C:E05	ES 147
	ES 147
M00039600A:A11	ES 148
	ES 148
	ES 148
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	ES 148
	ES 148
	ES 148
	ES 148
	ES 148
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Clone Name	The same of the sa
M00039626D:F04	ES 148
	ES 148
M00039629D:B04	ES 148
M00039630A:C08	ES 148
M00039630C:H04	ES 148
M00039641 A:A05	ES 148
M00039641C:D07	ES 148
M00039642D:B12 M00039642D:H09	ES 148 ES 148
M00039642D.H09 M00039643C:B04	ES 148
M00039645C:E01	ES 148
M00039647A:H11	ES 148
M00039047A.H11	ES 148
M00039740B:F10	ES 148
M00039740B:F10	ES 148
M00039755A:B08	ES 148
M00039755A.B08	ES 148
M00039760B:B08	ES 148
M00040131B:D11	ES 148
M00040131C:F03	ES 148
M00040131D:G08	ES 148
M00040133B:B03	ES 148
M00040136C:F08	ES 148
M00040138B:H03	ES 148
M00040141D:F05	ES 148
M00040143A:H05	ES 148
M00040145D:D03	ES 148
M00040147D:H11	ES 148
M00040160B:A10	ES 148
M00040162A:E01	ES 148
M00040169B:F08	ES 148
†	ES 148
M00040174C:E10	ES 148
M00040174D:G03	ES 148
M00040181B:H09	ES 148
	ES 148
	ES 148
M00040183A:F07	ES 148
M00040184C:A11	ES 148
M00040191A:B09	ES 148
M00040221A:G11	ES 148
	ES 148
	ES 148
	ES 148
M00040230A:H02	ES 148

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Clone Name:	Actibe
M00040231B:C08	ES 148
M00040232D:B07	ES 148
M000402322:B01	ES 148
M000402337::102 M00040233C:G05	ES 148
M00040253C:G05	ES 148
M00040252C:C00 M00040253C:A05	ES 148
M00040254B:C10	ES 148
M00040254B.C10	ES 148
	ES 148
M00040257D:H10	ES 148
M00040260B:D02	ES 148
M00040260C:D04	
M00040261C:F01	ES 148
M00040262B:B06	ES 148
M00040264D:G05	ES 148
M00040265D:B07	ES 148
M00040265D:C08	ES 148
M00040267A:E06	ES 148
M00040267C:C04	ES 148
M00040271B:E12	ES 148
M00040271C:D08	ES 148
M00040273B:H12	ES 148
M00040274A:D07	ES 148
M00040274A:H11	ES 148
M00040280C:H05	ES 148
M00040281D:B01	ES 148
M00040282A:A03	ES 148
M00040286C:C02	ES 148
M00040287C:B09	ES 148
M00040287D:D07	ES 148
M00039746C:A08	ES 148
M00039746C:G09	ES 148
M00039746C:H05	ES 148
	ES 148
M00039746D:D11	ES 148
	ES 148
	ES 148
	ES 148
M00039761D:E10	ES 148
M00039761B:E10	ES 148
	ES 148
M00039764C:D07	ES 148
	ES 149
	ES 149
M00039767B:A04 M00039767C:E12	ES 149
l	ES 149
M00039770A:G11	E3 147

Clone Name	"Tube"
M00039770C:E04	ES 149
M00039942D:C01	ES 149
M00039943B:F10	ES 149
M00039945C:F09	ES 149
M00039946B:F08	ES 149
M00039947A:D06	ES 149
M00039947C:G03	ES 149
M00039948A:E03	ES 149
M00039948D:D11	ES 149
M00039951A:B07	ES 149
M00039951B:B12	ES 149
M00039951B:C03	ES 149
M00039955C:C04	ES 149
M00039957C:C09	ES 149
M00039957D:A12	ES 149
M00039958A:A08	ES 149
M00039958C:B09	ES 149
M00040201C:G11	ES 149
M00040202A:F05	ES 149
M00040203A:H06	ES 149
M00040203B:A05	ES 149
M00040203D:H11	ES 149 ES 149
M00040206A:A07 M00040207B:D08	ES 149
M00040207B.D08	ES 149
M00040208A:C03	ES 149
M00040208D:G09	ES 149
M00040217D:B07	ES 149
M00040218C:C02	ES 149
M00040219B:D02	ES 149
M00040219D:E08	ES 149
M00040291D:C05	ES 149
M00040293D:G04	ES 149
M00040294D:D12	ES 149
M00040296D:E09	ES 149
M00040298B:G02	ES 149
M00040299B:F10	ES 149
M00040313C:D05	ES 149
M00040313D:E04	ES 149
M00040314D:H05	ES 149
M00040317A:H03	ES 149
M00040317D:F02	ES 149
M00040318A:B02	ES 149
	ES 149
M00040320D:F02	ES 149

Clone Name	≫Tube*
M00040323B:C12	ES 149
M00040323C:G11	ES 149
M00040326A:F04	ES 149
M00040327B:G06	ES 149
M00040332D:B05	ES 149
M00040333D:G05	ES 149
M00040334D:B02	ES 149
M00040334D:C07	ES 149
M00040342B:D12	ES 149
M00040345D:A09	ES 149
M00040346A:C11	ES 149
M00040347D:F09	ES 149
M00040349D:B09	ES 149
M00040351B:F02	ES 149
M00040351D:A11	ES 149
M00040364A:E05	ES 149
M00040366A:B01	ES 149
M00040368A:A12	ES 149
M00040368A:F01	ES 149
M00040368D:E09	ES 149
M00040371C:H05	ES 149
M00040375C:B06	ES 149
M00040376C:G02	ES 149
M00040377C:G07	ES 149
M00040383A:H02	ES 149
M00040383D:C04	ES 149
M00040385C:D02	ES 149
M00040386A:A02	ES 149
M00040387C:E07	ES 149
M00040387D:H05	ES 149
M00040390A:H02	ES 149
M00040390B:F02	ES 149
M00040391A:D10	ES 149
M00040392B:H01	ES 149
M00040392C:B12	ES 149
M00040394A:D04	ES 149
M00040395B:D11	ES 149
M00042534A:A05	ES 149
M00042538B:E06	ES 149
M00042543C:G04	ES 149
M00042558A:F03	ES 149
M00042560A:F12	ES 149
M00042565C:A08	ES 149
M00042566C:C05	ES 149
M00042567B:H10	ES 149
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Clone Name	Tube
M00042693D:E04	ES 149
M00042696B:E05	ES 149
M00042697D:C07	ES 150
M00042698D:D10	ES 150
M00042698D:E01	ES 150
M00042702B:G02	ES 150
M00042704A:F09	ES 150
M00042711B:A11	ES 150
M00042717A:C07	ES 150
M00042737C:H04	ES 150
M00042740A:E09	ES 150
M00042742D:D05	ES 150
M00042887C:D07	ES 150
M00042895A:D10	ES 150
M00042895C:G01	ES 150
M00042902D:B08	ES 150
M00042904B:E07	ES 150
M00042905A:F11	ES 150
M00042905B:C03	ES 150
M00042905D:D02	ES 150
M00042347D:H11	ES 150
M00042348B:E05	ES 150
M00042349D:D07	ES 150
M00042431B:G08	ES 150
M00042431C:F01	ES 150
M00042431D:C10	ES 150
M00042432D:E02	ES 150
M00042435A:A11	ES 150
M00042436B:H09	ES 150
M00042437A:D04	ES 150
M00042439B:B03	ES 150
M00042439B:D03	ES 150
M00042440B:E09	ES 150
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M00042515C:F08	ES 150
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M00042756B:F11	ES 150
M00042756D:A10	ES 150
M00042759B:G11	ES 150
M00042760A:C12	ES 150
M00042765C:D04	ES 150
M00042767B:G10	ES 150

Clone Name	Tube
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M00042770B:B12	ES 150
M00042770C:C04	ES 150
M00042771C:F06	ES 150
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	ES 150
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M00042790C:C07	ES 150
M00042792A:H01	ES 150
M00042797D:D10	ES 150
M00042799D:F08	ES 150
M00042800A:A03	ES 150
M00042802C:C04	ES 150
M00042806C:F07	ES 150
M00042807D:D05	ES 150
M00042823C:C02	ES 150
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M00042841D:H07	ES 150
M00042849D:F11	ES 150
M00042852B:A03	ES 150
M00042852C:A01	ES 150
M00042856B:H02	ES 150
M00042352C:H03	ES 150
M00042352D:C01	ES 150
M00042352D:G09	ES 150
M00042332D:G09	ES 150
M00042448C:H12	ES 150
M00042453B:G09	ES 150
M00042433B:G09	ES 150
M00042518D:A00	ES 150
M00042318D.D04 M00043296B:G09	ES 150
M00043296B:G09 M00043304B:D05	ES 150
MUUU43304B:DU3	ES 150
M00043304C:D02	ES 150
M00043305B:G02	
M00043306C:B03	ES 150
M00043306D:B07	ES 150
M00043310C:G06	ES 150
M00043311C:E03	ES 150
M00043312C:E08	ES 150
M00043320B:A07	ES 150
M00043324D:H11	ES 150
M00043328D:H02	ES 150
M00043332C:G04	ES 150
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Clone Name	Tube
M00043334B:A10	ES 150
M00043338B:A03	ES 150
M00043338B:C11	ES 150
M00043339A:F11	ES 150
M00043340B:H08	ES 150
M00043344D:E04	ES 150
M00043345C:A06	ES 150
M00043346A:G01	ES 150
M00043350D:B11	ES 151
M00043351D:A11	ES 151
M00043352D:B05	ES 151
M00043352D:C03	ES 151
M00043359B:D10	ES 151
M00043359C:G01	ES 151
M00043361B:A01	ES 151
M00043366A:A02	ES 151
M00043366C:H05	ES 151
M00043367B:A08	ES 151
M00043368C:F09	ES 151
M00043370B:C08	ES 151 ES 151
M00043372C:G05	
M00043377A:C03	ES 151 ES 151
M00043378A:H10 M00043379D:H02	ES 151
M00043379D:1102 M00043383C:F12	ES 151
M00043383D:A02	ES 151
M00043384B:B02	ES 151
M00043386A:B08	ES 151
M00043389C:E03	ES 151
M00043389D:D07	ES 151
M00043391A:C10	ES 151
M00043391A:G08	ES 151
M00043392D:C11	ES 151
M00043393A:B08	ES 151
M00043401D:G08	ES 151
M00043402C:D08	ES 151
M00043405A:D11	ES 151
M00043405C:G12	ES 151
M00043405C:G02	ES 151
M00043406B:G12	ES 151
M00043407C:E05	ES 151
M00043408B:D11	ES 151
M00043409B:B03	ES 151 ES 151
M00043410C:A09	120 121
M00043411B:D08	ES 151

Clone! Name	Tube
M00043411D:H06	ES 151
M00042584B:C10	ES 151
M00042623D:D07	ES 151
M00042625C:B04	ES 151
M00042626B:D08	ES 151
M00042627C:D01	ES 151
M00042630A:C05	ES 151
M00042955C:D05	ES 151
M00042956C:B06	ES 151
M00042960D:H08	ES 151
M00042962D:C05	ES 151
M00042964D:A03	ES 151
M00042966B:F07	ES 151
M00042966C:E06	ES 151
M00042970C:A04	ES 151
M00042970C:H10	ES 151
M00042976A:H04	ES 151
M00042979B:E02	ES 151
M00042981B:D11	ES 151
M00042983C:A11	ES 151
M00042983C:G06	ES 151
M00042986C:G12	ES 151
M00042988A:F06	ES 151
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M00042998A:G04	ES 151
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M00043003C:D08	ES 151
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M00043015A:H10	ES 151
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M00043026C:D07	ES 151
M00043028A:G05	ES 151
M00043029C:A06	ES 151
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M00043034D:C01	ES 151
M00043036C:E05	ES 151
M00043036D:C09	ES 151
M00043040B:B07	ES 151
M00043044B:A12	ES 151
M00043044D:A09	ES 151
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M00043046D:B11	ES 151

Glone Name	Tube
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M00043066B:H11	ES 151
M00043067D:D10	ES 151
M00043125A:B11	ES 151
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M00042614B:B05	ES 151
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M00043081D:F05	ES 151
M00043087B:G07	ES 151
M00043093C:G11	ES 151
M00043095A:F09	ES 152
M00043096A:G04	ES 152
M00043108A:F06	ES 152
M00043109C:G01	ES 152
M00043131B:A09	ES 152
M00043133B:C11	ES 152
M00043138D:B11	ES 152
M00043143B:A10	ES 152
M00043148C:A09	ES 152
M00043154A:B07	ES 152
M00043162A:B08	ES 152
M00043162D:C12	ES 152
M00043164C:E12	ES 152
M00043165B:G01	ES 152
M00043173D:G03	ES 152
M00043184A:H08	ES 152
M00043187A:C04	ES 152
M00043191A:A07	ES 152
M00043192C:B12	ES 152
M00043200A:H09	ES 152
M00043200B:C08	ES 152
M00043202B:F01	ES 152
M00043203A:B09	ES 152
M00043210C:E05	ES 152
M00043211A:F01	ES 152
M00043213B:B12	ES 152
M00043215A:D02	ES 152
M00043220B:C04	ES 152
M00042591D:H03	ES 152
M00042592A:H10	ES 152
M00042593A:C02	ES 152
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M00042593C:G06	ES 152
M00042595A:A11	ES 152
M00042595A:B01	ES 152
M00042596B:F06	ES 152
M00042596C:D07	ES 152
M00042597B:E12	ES 152
M00042377B:E12	ES 152
M00043417C:D05	ES 152
M00043417C:B03	ES 152
M00043419D:A10	ES 152
M00043419D:A10	ES 152
M00043428D:G08	
M00043431D:B08	ES 152
M00043433B:G09	ES 152
M00043433C:G07	ES 152
M00043437D:D04	ES 152
M00043440C:B07	ES 152
M00043446C:E12	ES 152
M00043447A:C07	ES 152
M00043449A:E12	ES 152
M00043450C:C06	ES 152
M00043453B:B09	ES 152
M00043458A:B12	ES 152
M00043461D:C02	ES 152
M00043461D:E06	ES 152
M00043465B:H02	ES 152
M00043465C:A03	ES 152
M00043465C:C09	ES 152
M00043476A:F07	ES 152
M00043483B:G10	ES 152
M00043491C:F04	ES 152
M00043492A:E01	ES 152
M00043513D:G08	ES 152
M00043516B:H09	ES 152
M00043518B:D06	ES 152
M00043526B:D10	ES 152
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M00043528C:A02	ES 152
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€ Glone Name	Tube
M00043638A:D06	ES 152
M00043640C:E03	ES 152
M00043648A:G07	ES 152
M00043649B:E07	ES 152
M00001338C:B02	ES 153
M00001338C:F05	ES 153
M00001338D:D01	ES 153
M00001340D:F07	ES 153
M00001344D:E08	ES 153
M00001346B:G11	ES 153
M00001348B:B03	ES 153
M00001349C:B04	ES 153
M00001351B:E11	ES 153
M00001352B:B02	ES 153
M00001353A:H07	ES 153
M00001353C:A05	ES 153
M00001353D:E05	ES 153
M00001356D:E06	ES 153
M00001358A:E08	ES 153
M00001359A:H10	ES 153
M00001361A:C12	ES 153
M00001361B:A12	ES 153
M00001362A:F09	ES 153
M00001364A:C09	ES 153
M00001364C:H10	ES 153
M00001368A:A08	ES 153
M00001368A:B07	ES 153
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M00001369A:G06	ES 153
M00001374A:B02	ES 153
M00001374C:B10	ES 153
M00001375B:D04	ES 153
M00001378C:E10	ES 153
M00001379A:F09 M00001382D:A07	ES 153
M00001382D:H08	ES 153 ES 153
M00001382D:H08	
M00001384A:A07 M00001385A:E07	ES 153 ES 153
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M00001393B:C03	ES 153
M00001393C:E08	ES 153
M00001393C:F04	ES 153
M00001393D:E02	ES 153
M00001396B:B01	ES 153
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M00001396D:H02	ES 153
M00001397C:H08	ES 153
M000013978:B01	ES 153
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M00001399C:A01	ES 153
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M00001406B:H09	ES 153
M00001406D:F06	ES 153
M00001410A:G10	ES 153
M00001416B:A05	ES 153
M00001421B:E07	ES 153
M00001422B:D06	ES 153
M00001424B:H06	ES 153
M00001424D:D02	ES 153
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M00001428B:C10	ES 153
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M00001433B:E02	ES 153
M00001442A:F08	ES 153
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M00001449B:H10	ES 153
M00001451C:E10	ES 153
M00001460C:E10	ES 153
M00001461D:B10	ES 153
M00001461D:C10	ES 153
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M00001466B:F03	ES 153
M00001467C:D04	ES 153
M00001477D:G09	ES 153
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Clone Name	Tube
M00001518B:D10	ES 153
M00001528C:C03	ES 153
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M00001533C:G11	ES 153
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M00001534C:E07	ES 153
M00001535B:B10	ES 153
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M00001537B:H10	ES 153
M00001538B:A07	ES 153
M00001539C:F12	ES 154
M00001542B:F09	ES 154
M00001543C:A08	ES 154
M00001544B:B05	ES 154
M00001544B:E06	ES 154
M00001546B:C11	ES 154
M00001548B:D06	ES 154
M00001550A:H06	ES 154
M00001550D:B11	ES 154
M00001551D:D01	ES 154
M00001551D:H09	ES 154
M00001554C:G10	ES 154
M00001558A:E06	ES 154
M00001559A:H09	ES 154
M00001561D:H04	ES 154
M00001562B:B02 M00001562D:B07	ES 154
M00001565A:H05	ES 154 ES 154
M00001568C:A03	ES 154
M00001570A:B07	ES 154
M00001591B:H05	ES 154
M00001596A:D02	ES 154
M00001600B:G01	ES 154
M00001605B:B05	ES 154
M00001606B:A10	ES 154
M00001606D:D06	ES 154
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M00001607D:H09	ES 154
M00001609D:C11	ES 154
M00001616D:F03	ES 154
M00001617C:F10	ES 154
M00001618C:D01	ES 154
M00001619C:H09	ES 154
M00001620B:A03	ES 154
M00001623D:A10	ES 154

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M00001623D:E12	ES 154
M00001624A:C01	ES 154
M00001625D:B04	ES 154
M00001626A:D07	ES 154
M00001632C:A10	ES 154
M00001633D:C11	ES 154
M00001637D:C12	ES 154
M00001648A:D10	ES 154
M00001661D:F06	ES 154
M00001663A:A12	ES 154
M00001671A:H10	ES 154
M00001671C:F03	ES 154
M00001675B:D06	ES 154
M00001677B:H08	ES 154
M00001677B:1108	ES 154
M00001683B:F11	ES 154
M00001684D:E04	ES 154
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M00003753A:C11	ES 154
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M00003763B:B10	ES 154
M00003764B:F11	ES 154
M00003764B:H11	ES 154
M00003764D:F07	ES 154
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M00003771D:A03	ES 154
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M00003780A:G01	ES 154
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M00003785D:F07	ES 154
M00003787D:A10	ES 154
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M00003808B:E07	ES 154

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M00003812C:A03	ES 154
M00003814A:G05	ES 154
M00003819B:B01	ES 154
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M00003822C:A09	ES 154
M00003822D:A02	ES 154
M00003823B:A06	ES 154
M00003825A:H10	ES 154
M00003828A:D11	ES 154
M00003830B:C06	ES 154
M00003830C:D02	ES 154
M00003837C:D10	ES 154
M00003839C:H10	ES 154
M00003842D:D11	ES 154
M00003842D:H09	ES 154
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M00003851B:A01	ES 155
M00003854B:F07	ES 155
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M00003915C:D10	ES 155
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M00003965D:D11	ES 155
M00003968C:G03	ES 155

Clone Name	Tube
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M00003972C:F07	ES 155
M00003974C:E11	ES 155
M00003974D:E02	ES 155
M00003979B:A04	ES 155
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M00004085B:H02	ES 155
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M00004096D:F02	ES 155
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M00004103C:E10	ES 155
M00004104A:A12	ES 155
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M00004118C:D12	ES 155

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M00004122C:D01	ES 155
M00004134A:A08	ES 155
M00004136C:B12	ES 155
M00004139B:F01	ES 155
M00004141A:D01	ES 155
M00004141B:B01	ES 155
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M00004143B:B04	ES 155
M00004144D:B02	ES 155
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M00004146B:E08	ES 155
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M00004158B:E03	ES 155
M00004158D:E08	ES 155
M00004159C:D10	ES 155
M00004159D:F12	ES 155
M00004160D:F06	ES 155
M00004160D:G05	ES 155
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	ES 156
M00004172D:B12	ES 156
M00004172D:F04	ES 156
M00004175D:E06	ES 156
M00004176C:A09	ES 156
M00004179C:B06	ES 156
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M00004192C:B06	ES 156
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Clone Name	Mube
M00004204C:H08	ES 156
M00004207C:A04	ES 156
M00004208A:D08	ES 156
M00004210A:A03	ES 156
M00004212D:C03	ES 156
M00004214A:E05	ES 156
M00004214D:A05	ES 156
M00004215B:C05	ES 156
M00004220D:C11	ES 156
M00004225D:E03	ES 156
M00004229B:B06	ES 156
M00004230D:B05	ES 156
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M00004303C:C03	ES 156
M00004316A:B03	ES 156
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M00004320C:E07	ES 156
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M00004330A:A01	29 1 20

Clone Name : 1	Tübe
M00004336A:A01	ES 156
M00004341C:A09	ES 156
M00004341C:E05	ES 156
M00004344A:G11	ES 156
M00004344D:C12	ES 156
M00004347B:E04	ES 156
M00004347C:A05	ES 156
M00004350A:A04	ES 156
M00004351B:G07	ES 156
M00004352A:D08	ES 156
M00004357B:B06	ES 156
M00004358B:G02	ES 156
M00004359A:E01	ES 156
M00004360C:D09	ES 156
M00004365C:C09	ES 156
M00004365C:G11	ES 156
M00004366D:C11	ES 156
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M00004372A:E12	ES 156
M00004376D:A12	ES 156
M00004385C:H12	ES 156
M00004393C:D06	ES 156
M00004406A:G09	ES 156
M00004416B:G10	ES 156
M00004418B:A11	ES 156
M00004419A:G02	ES 156
M00004420D:E05	ES 156
M00004430A:A05	ES 156
M00004430B:B10	ES 157
M00004443C:F07	ES 157
M00004462D:D12	ES 157
M00004502A:D12	ES 157
M00004507D:E03	ES 157
M00004509B:B10 M00004509D:C06	ES 157
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M00004603B:E02	ES 157
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M00004608A:C10	ES 157
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M00004608A:H04 M00004609A:E09	ES 157
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M00023394D.D10	ES 157
M00026809A:H08	ES 157
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M00026836B:H03	ES 157
M00026842B:A01	ES 157
M00026842D:C02	ES 157
M00026850B:C09	ES 157
M00026856B:G03	ES 157
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M00027057C:D10	ES 157
M00027064B:D06	ES 157
M00027081A:A08	ES 157
M00027093 A:H02	ES 157
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M00027178B:A11	ES 157
M00027182B:G06	ES 157
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M00027215B:B12	ES 157
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M00027262A:A07	ES 157
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M00027316C:C03	ES 157

Glone Name	Tube
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M00027319D:F07	ES 157
M00027332B:H09	ES 157
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M00027457B:E11	ES 157
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M00027475D:A01	ES 157
M00027480C:E09	ES 157
M00027485C:F07	ES 157
M00027506B:G01	ES 157
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M00027527B:C05	ES 157
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M00027724D:D04	ES 157
M00027734D:C03	ES 157
M00027746A:D06	ES 157
M00027801B:D07 M00027806C:H05	ES 157
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M00028784A:D12	ES 158
M00028786B:A04	ES 158
M00031370B:C01	ES 158
M00031416D:H05	ES 158
M00031484A:D03	ES 158
M00031485B:G05	ES 158
M00032471D:A05	ES 158
M00032473B:A03	ES 158
M00032474A:G03	ES 158
M00032475A:A06	ES 158
M00032489B:G12	ES 158
M00032490D:E08	ES 158
M00032494C:H08	ES 158
M00032497D:B10	ES 158
M00032504B:B10	ES 158
M00032507D:G08	ES 158
M00032508A:E03	ES 158
M00032505A:E03	ES 158
M00032517C:E10	ES 158
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M00032534B:E12	ES 158
M00032534B.E12	ES 158
M00032541C:G03	
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M00032556D:A03	1
M00032562C:F01	ES 158
M00032567B:G05	ES 158
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M00032580D:A09	ES 158
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M00032584A:D06	ES 158
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M00032614C:B10	ES 158
M00032614D:D08	ES 158
M00032620B:F06	ES 158
M00032621A:F11	ES 158
M00032628C:B06	ES 158
M00032634B:D09	ES 158
M00032637A:F09	ES 158
M00032638B:F02	ES 158
M00032644C:B05	ES 158
M00032645D:C01	ES 158
M00032647B:F06	ES 158
M00032652C:C07	ES 158
M00032666A:C02	ES 158
M00032671B:D06	ES 158
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M00032676C:C10	ES 158
M00032688C:A03	ES 158
M00032700A:E09	ES 158
M00032707D:F08	ES 158
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M00032723D:H02	ES 158
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M00032734B:E12	ES 158
M00032734C:C05	ES 158
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M00032753A:C07	ES 158
M00032759A:A03	ES 158
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Clone Name	Jubes
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M00032829D:A05	ES 159
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M00032831A:E09	ES 159
M00032835D:G04	ES 159
M00032836B:A07	ES 159
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M00032908A:D08	ES 159
M00032918D:B04	ES 159
M00032928C:D02	ES 159
M00032944A:B07	ES 159
M00032945D:B07	ES 159
M00032979D:C11	ES 159
M00032979D:H07	ES 159
M00032985D:G09	ES 159
M00032987B:F01	ES 159
M00032988B:G01	ES 159
M00033006A:F10	ES 159
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M00033028D:C10	ES 159
M00033037B:F04	ES 159
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M00033081D:D11	ES 159
M00033144A:D02	ES 159
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M00033149B·E10	ES 159
M00033150B:E02	ES 159
M00033150C:A11	ES 159
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M00033246A:H12	ES 159
M00033248D:H11 M00033264B:E06	ES 159
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M00033303C:F09	ES 159
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M00033359C:H05	ES 159
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M00038619B:F09	ES 159
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M00039125D:H12	ES 160
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M00039140A:F05	ES 160
M00039143A:F04	ES 160
M00039143D:C10	ES 160
M00039146B:G04	ES 160
M00039140D:C04	ES 160
M00039165D:C04	ES 160
M00039103D.C04 M00039175A:F01	ES 160
M00039173A:F01	ES 160
M00039207A:F07	ES 160
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M00039223A:D11	ES 160
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M00039251C:H12	ES 160
M00039251D:B08	ES 160
M00039255D:B01	ES 160
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M00039270D:D02	ES 160
M00039275B:E02	ES 160
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M00039344C:A11	ES 160
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M00039440C:G06	ES 160
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M00039668C:F01	ES 160
M00039672C:D05	ES 160
M00039686C:C01	ES 160
M00039694C:H01	ES 160
M00039698C:B03	ES 160
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M00039710B:E01	ES 160
M00039785C:H12	ES 160
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M00039806B:D05	ES 160 ES 160
M00039820B:F06 M00039822A:H02	ES 160
M00039826B:F09	ES 160
M00039826D:E04	ES 160
M00039828B:H06	ES 160
M00039829B:E01	ES 160
M00039860B:E01	ES 160
M00039860D:B02	ES 160
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M00039915B:E08	ES 160
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M00004832D:G04	ES 160
M00004836B:C02	ES 161
M00004839B:C12	ES 161
M00004843A:G12	ES 161
M00004846A:A10	ES 161
M00004850A:B02	ES 161
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M00004856D:F09	ES 161
M00004873B:G04	ES 161
M00004876B:A06	ES 161
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M00005014B:F02	ES 161
M00005016C:E04	ES 161
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M00005422B	:B08	ES	161	_
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M00005434A			161	
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M00005452B		ES		
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M00005512B			161	
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M00005606D			61	
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M00005622A:H02	ES 161
M00005623B:G01	ES 161
M00005626D:G11	ES 161
M00005634A:F07	ES 161
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M00005703D:G10	ES 161
M00005710B:H03	ES 162
M00005743D:A12	ES 162
M00005763D:A01	ES 162
M00005766D:D12	ES 162
M00005771D:C02	ES 162
M00005819D:F09	ES 162
M00005822C:A04	ES 162
M00006576D:C02	ES 162
M00006577A:H10	ES 162
M00006582D:A09	ES 162
M00006585A:D07	ES 162
M00006585A:F09	ES 162
M00006586D:D04	ES 162
M00006592A:A12	ES 162
M00006595B:C10	ES 162
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M00006631D:D02	ES 162
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M00006646A:A07	ES 162
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M00006714C:D06	ES 162
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M00006739B:A04	ES 162
M00006740B:A09	ES 162
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M00006743A:H11	ES 162
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M00006756C:A02	ES 162

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M00007048C:A12	ES 162
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M00007126A:A02	ES 162
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M00007154A:E06	ES 162
M00007155C:D07	ES 162
M00007155D:C09	ES 162
M00007158D:D03	ES 162
M00007178A:C02	ES 162
M00007195C:E11	ES 162
M00007197B:B05	ES 162
M00007202B:F01	ES 162
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M00007969D:C01	ES 162
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M00007975C:A10	ES 162
M00007975D:F12	ES 162
M00007980A:B01	ES 162
M00007980B:A07	ES 162
M00007981C:F07	ES 162
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M00008016B:E09	ES 162
M00008019B:A01	ES 162
M00008020D:D05	ES 162
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M00008021C:G12	ES 162
M00008045C:A05	ES 162
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M00008065D:A07	ES 163
M00008071D:H03	ES 163
M00008073A:D01	ES 163
M00008073D:D01	ES 163
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M00021678D:H04	ES 163
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M00021697C:B07	ES 163
M00021700D:H03	ES 163
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M00021958B:E08	ES 163

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M00021967D:H06	ES 163
M00021971C:B11	ES 163
M00021974D:F01	ES 163
M00021981A:C02	ES 163
M00021991D:F09	ES 163
M00021998B:D09	ES 163
M00022009C:A08	ES 163
M00022016B:F01	ES 163
M00022032A:G05	ES 163
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M00022096D:A03	ES 163
M00022103C:D05	ES 163
M00022104A:G08	ES 163
M00022117C:A02	ES 163
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M00022140D:A07	ES 163
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M00022170C:C01	ES 163 ES 163
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M00022103A:B03	ES 163
M00022193C:C09	ES 163
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M00022202C:C04	ES 163
M00022208B:D03	ES 163
M00022208C:E04	ES 163
M00022208C:F08	ES 163
M00022212D:G02	ES 163
M00022216D:D10	ES 163
M00022218B:B12	ES 163
M00022220A:A07	ES 163
M00022224A:C07	ES 163
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M00022228B:B11	ES 163

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M00022237D:D06	ES 163
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M00022240D:B11	ES 163
M00022249D:C01	ES 163
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M00022262A:F06	ES 163
M00022262B:B06	ES 163
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M00022363C:D05	ES 163
M00022367D:G11	ES 163
M00022368A:B11	ES 163
M00022372D:H12	ES 163
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Clone Name	LATABA
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M00022494D:A05	ES 164
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M00022516B:C05	ES 164
M00022525B:D09	ES 164
M00022530B:C04	ES 164
M00022537B:C06	ES 164
M00022546B:E05	ES 164
M00022559D:G10	ES 164
M00022563B:C08	ES 164
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M00022618C:E04	ES 164
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M00022648A:D08	ES 164
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M00022662D:H03	ES 164
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M00040034A:E06	ES 164
M00040034R:G02	ES 164
M00040034B:G02	ES 164
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M00040043B:1107	ES 164
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M00040094B:C08	ES 164 ES 164
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M00022738D:G08	ES 164
M00022740C:H11	ES 165
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M00022835C:A09	ES 165
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	ES 165
M00022904C:D04	ES 165
M00022924B:A05	ES 165

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M00022945B:F11	ES 165
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M00022953B:D06	ES 165
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M00023007D:D03	ES 165
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M00039537A:F08	ES 165
M00039564D:D04	ES 165
M00039594C:B06	ES 165
M00039598A:E04	ES 165
M00039630D:B07	ES 165
M00039642A:A08	ES 165
M00039642C:F08	ES 165
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M00039647A:A02	ES 165
M00039647B:A02	ES 165
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M00040162A:E02	ES 165
M00040169A:G06	ES 165
M00040173D:A04	ES 165
M00040174D:G06	ES 165
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M00040252C:G05	ES 165
M00040267D:A12	ES 165
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M00040289D:C06	ES 165
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M00040349D:D07	ES 165
M00040351A:C08	ES 165
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M00040367A:C08	ES 165
M00040381A:B06	ES 165
M00040384B:E04	ES 165
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M00042525B:H01	ES 165
M00042528C:H01	ES 165
M00042554A:D01	ES 165
M00042557D:B06	ES 165
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M00042579A:D09	ES 165
M00042719A:G08	ES 165
M00042722C:C09	ES 165
M00042724A:G06	ES 165
M00042732B:H06	ES 165
M00042734A:F05	ES 165
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M00042894C.A11	ES 165
M00042915B:G11	ES 165
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M00055456C:H06	ES 166
M00055466A:F06	ES 166
M00055468A:A08	ES 166
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M00055639A:E06	ES 166
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M00055676A:G02	ES 166
M00055724B:E04	ES 166
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M00055925D:B07	ES 166
M00055961C:B10	ES 166
M00055975B:F09	ES 166
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M00056004B:C05	ES 166
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M00056133A:E11	ES 166 ES 166
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M00056217D:E10	ES 166
M00056220D:G02	ES 166
M00056230D:E07	ES 166
<u></u>	ES 166
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M00056320B:A03	ES 166
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M00056345D:A04	ES 166
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M00056458C:E01	ES 166
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M00042433A:E11	ES 166
M00042462B:C02	ES 166
	ES 166
M00042766C:D05	ES 166

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Clone Name	畫Tube:
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M00042822A:H04	ES 166
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M00042860B:C07	ES 166
M00042863D:F09	ES 166
M00042878D:F05	ES 166
M00042878D:G06	ES 166
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M00042352D:B03	ES 166
M00042449B:F05	ES 166
M00042457C:B06	ES 166
M00042516B:D01	ES 166
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M00043313D:E09	ES 166
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M00043336D:B03	ES 166
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M00054683D:G11	ES 166
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Glone Name:	a lube
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M00054781D:A11	ES 167
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M00054817D:A11	ES 167
M00054818B:F10	ES 167
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M00054866B:C08	ES 167
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M00054931D:E10	ES 167
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M00055002B:E08	ES 167
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M00055023A:E11	ES 167
M00055043B:H08	ES 167
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M00055239D:F11	ES 167
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	ES 167
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M00042960B:C06	ES 167
M00042967D:C01	ES 167
M00042970C:B01	ES 167
M00042972C:F04	ES 167
M00042976D:C01	ES 167
M00042982D:A10	ES 167
M00042986D:E03	ES 167
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M00043015D:D05	ES 167
M00043016B:F09	ES 167
M00043017C:D08	ES 167
M00043063C:H05	ES 167
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M00043113C:G09	ES 167

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M00043077C:D12	ES 167
M00043077C:G10	ES 167
M00043099A:H04	ES 167
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M00043465C:H11	ES 167
M00043470A:C10	ES 167
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CLAIMS

We claim:

- 1. A library of polynucleotides, the library comprising the sequence information of at least one of SEQ ID NO:1-3351.
- 2. The library of claim 1, wherein the library is provided on a nucleic acid array.
- 3. The library of claim 1, wherein the library is provided in a computer-readable format.
- 4. The library of claim 1, wherein the library comprises a polynucleotide corresponding to a gene differentially expressed in a cancer cell of high metastatic potential relative to a control cell, wherein the control cell is a normal cell or a cell of low metastatic potential, wherein the expression is greater in the metastatic tissue, and wherein the sequence is selected from the group consisting of SEQ 1D NOs:14, 137, 151, 152, 171, 200, 254, 262, 271, 348, 412, 472, 507, 520, 530, 588, 623, 637, 660, 678, 680, 700, 714, 774, 812, 834, 901, 937, 976, 1168, 1333, 1352, 1520, 1524, 1546, 1550, 1574, 1580, 1590, 1599, 1607, 1622, 1706, 1752, 1768, 1769, 1780, 1781, 1799, 1803, 1811, 1851, 1856, 1867, 1872, 1875, 1884, 1919, 1923, 1939, 1975, 2024, 2045, 2060, 2071, 2118, 2119, 2128, 2135, 2177, 2181, 2184, 2185, 2190, 2193, 2232, 2239, 2283, 2311, 2314, 2338, 2378, 2393, 2394, 2395, 2398, 2460, 2490, 2505, 2514, 2540, 2542, 2597, 2607, 2640, 2657, 2669, 2670, 2674, 2679, 2684, 2707, 2724, 2757, 2776, 2804, 2818, 2906, 2959, 2964, 2968, 2976, 2980, 2987, 3010, 3043, 3047, 3050, 3071, 3072, 3092, 3095, 3097, 3140, 3157, 3173, 3187, 3203, 3210, 3212, 3220, 3236, 3249, 3264, 3284, 3288, 3305, 3309, 3318, 3330, 3331, and 3335.
- 5. The library of claim 1, wherein the library comprises a polynucleotide corresponding to a gene differentially expressed in normal colon tissue relative to colon cancer tissue, wherein the expression is greater in the cancer tissue, and wherein the sequence is selected from the group consisting of SEQ 1D NOs:7, 164, 734, 836, 928, 965, 987, 1026, 1044, 1119, 1226, 1227, 1251, 1316, 1429, 1442, 1540, 1553, 1560, 1577, 1588, 1610, 1620, 1626, 1673, 2416, 2749, 2976, 3129 and 3132.

6. The library of claim 1, wherein the library comprises a polynucleotide corresponding to a gene differentially expressed in normal colon tissue relative to colon cancer tissue, wherein the expression is greater in normal tissue than cancer tissue, and wherein the sequence is selected from the group consisting of SEQ ID NOs:105, 198, 465, 489, 745, 859, 976, 1011, 1045, 1138, 1226, 1251, 1253, 1392, 1474, 1559, 1571, 1589, 1591, 1607, 1608, 1643, 1753, 1764, 1766, 1782, 1811, 2749, 2784, 2790, 2805, 2976, 3128, 3129, 3146, 3150, and 3151.

- 7. The library of claim 1, wherein the library comprises a polynucleotide corresponding to a gene differentially expressed in normal human prostate cells relative to human prostate cancer cells, wherein the expression is greater in normal cells than cancer cells, and wherein the sequence is selected from the group consisting of SEQ ID NOs:53, 446, 1410, 1754, 1801, 1845, 2060, 2143, 2632, 2899, and 3338.
- 8. The library of claim 1, wherein the library comprises a polynucleotide corresponding to a gene differentially expressed in normal human prostate cells relative to human prostate cancer cells, wherein the expression is greater in cancer cells than normal cells, and wherein the sequence is selected from the group consisting of SEQ ID NOs:86, 93, 687, 1269, 1581, 1647, 1649, 1710, 1717, 1772, 1960, 2987, 3128, 3132, 3150, 3222, and 3268.
- 9. An isolated polynucleotide comprising a nucleotide sequence having at least 90% sequence identity to an identifying sequence of SEQ 1D NOs:1-3351 or a degenerate variant or fragment thereof.
 - 10. A recombinant host cell containing the polynucleotide of claim 9.
 - 11. An isolated polypeptide encoded by the polynucleotide of claim 9.
 - 12. An antibody that specifically binds a polypeptide of claim 11.
 - 13. A vector comprising the polynucleotide of claim 9.
- 14. A method of detecting differentially expressed genes correlated with a cancerous state of a mammalian cell, the method comprising the step of:

detecting at least one differentially expressed gene product in a test sample derived from a cell suspected of being cancerous, wherein the gene product is encoded by a

gene corresponding to a sequence of at least one of SEQ ID NOs: 14, 137, 151, 152, 171, 200, 254, 262, 271, 348, 412, 472, 507, 520, 530, 588, 623, 637, 660, 678, 680, 700, 714, 774, 812, 834, 901, 937, 976, 1168, 1333, 1352, 1520, 1524, 1546, 1550, 1574, 1580, 1590, 1599, 1607, 1622, 1706, 1752, 1768, 1769, 1780, 1781, 1799, 1803, 1811, 1851, 1856, 1867, 1872, 1875, 1884, 1919, 1923, 1939, 1975, 2024, 2045, 2060, 2071, 2118, 2119, 2128, 2135, 2177, 2181, 2184, 2185, 2190, 2193, 2232, 2239, 2283, 2311, 2314, 2338, 2378, 2393, 2394, 2395, 2398, 2460, 2490, 2505, 2514, 2540, 2542, 2597, 2607, 2640, 2657, 2669, 2670, 2674, 2679, 2684, 2707, 2724, 2757, 2776, 2804, 2818, 2906, 2959, 2964, 2968, 2976, 2980, 2987, 3010, 3043, 3047, 3050, 3071, 3072, 3092, 3095, 3097, 3140, 3157, 3173, 3187, 3203, 3210, 3212, 3220, 3236, 3249, 3264, 3284, 3288, 3305, 3309, 3318, 3330, 3331, and 3335.

wherein detection of the differentially expressed gene product is correlated with a cancerous state of the cell from which the test sample was derived.

15. A method of detecting differentially expressed genes correlated with a cancerous state of a mammalian cell, the method comprising the step of:

detecting at least one differentially expressed gene product in a test sample derived from a cell suspected of being cancerous, wherein the gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOs:7, 164, 734, 836, 928, 965, 987, 1026, 1044, 1119, 1226, 1227, 1251, 1316, 1429, 1442, 1540, 1553, 1560, 1577, 1588, 1610, 1620, 1626, 1673, 1960, 2416, 2749, 2976, 2987, 3128, 3129, 3132, 3150, 3222, and 3268.

wherein detection of the differentially expressed gene product is correlated with a cancerous state of the cell from which the test sample was derived.